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#### (57) Abstract

The present invention relates generally to three novel human genes with gene regulatory function. These genes encode a zinc finger protein, a guanine nucleotide exchange protein and a heat shock protein or heat shock binding protein. The invention includes derivatives and mammalian animal, insect, nematodes, avian and microbial homologues of these genes. The present invention further provides pharmaceutical compositions and diagnostic agents as well as genetic molecules useful in gene replacement therapy and recombinant molecules useful in protein replacement therapy.



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THREE NOVEL GENES ENCODING A ZINC FINGER PROTEIN, A GUANINE, NUCLEOTIDE EXCHANGE FACTOR AND A HEAT SHOCK PROTEIN OR HEAT SHOCK BINDING PROTEIN

#### FIELD OF THE INVENTION

5 The present invention relates generally to a novel human gene and its derivatives and to mammalian, animal, insect, nematodes, avian and microbial homologues thereof. The present invention further provides pharmaceutical compositions and diagnostic agents as well as genetic molecules useful in gene replacement therapy and recombinant molecules useful in protein replacement therapy.

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#### **BACKGROUND OF THE INVENTION**

Bibliographic details of the publications referred to by author in this specification are collected at the end of the description.

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The increasing sophistication of recombinant DNA technology is greatly facilitating research and development in the medical and allied health fields. There is growing need to develop recombinant and genetic molecules for use in diagnosis and in conventional pharmaceutical preparations as well as in gene and protein replacement therapies.

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In work leading up to the present invention, the inventors sought to identify and clone human genes which might be useful as potential diagnostic and/or therapeutic agents. Molecules of particular interest targeted by the inventors were gene regulators including regulatory proteins, signal transducters and heat shock proteins.

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Gene expression generally requires interaction between a regulatory protein and an appropriate recognition sequence of a target gene. Regulatory proteins comprise in many cases a domain or motif which facilitates binding to DNA. One particular motif comprises small sequence units repeated in tandem with each unit folded about a zinc atom to form separate structural domains.

30 This motif is now referred to as a zinc finger domain. Such a domain is generally defined by the number of cysteine (C) and histidine (H) residues.

In addition, knowledge of cellular interaction in the control of cell proliferation is essential in the rational design of specific therapeutic strategies aimed at controlling proliferative disorders. Such proliferative disorders including a range of cancers, inflammatory conditions and atherosclerosis. An important aspect of cellular interaction is in signal transduction *via* receptors to intracellular transducers. One key signal transducer is Ras which couples the receptors for diverse extracellular signals to different effectors. Ras directly activates the downstream kinase Raf which in turn induces the mitogen activated protein kinase (MAPK) cascade.

Another regulatory mechanism involves heat shock proteins. The *Escherichia coli* heat shock protein, DnaJ, is the founding member of a family of proteins which are associated with protein folding, protein complex assembly and transit through subcellular components.

Prokaryotic and eukaryotic DnaJ homologues have a modular organisation consisting of a J domain, a glycine-rich spacer, CXXCXGXG [SEQ ID NO:1] repeats and a C-terminal region with no obvious sequence features, as well as additional sequences for protein targeting. The J domain is anticipated to mediate interaction with heat shock 70 proteins (Hsp70) and consists of some 70 amino acids, frequently located at the N-terminus of the protein.

In accordance with the present invention, a genes have been identified from the human genome which encodes proteins having a regulatory role. One gene, in accordance with the present invention encodes a protein with an N-terminal region resembling a zinc-finger domain of a novel type. Another gene encodes a protein involved in guanine nucleotide exchange factor (GEF) signalling pathways. Yet another gene encodes a protein which is a heat shock protein or heat shock-like protein which may have a role in tumour suppression.

#### SUMMARY OF THE INVENTION

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Throughout this specification, unless the context requires otherwise, the word "comprise", or variations such as "comprises" or "comprising", will be understood to imply the inclusion of a stated element or integer or group of elements or integers but not the exclusion of any other element or integer or group of elements or integers.

Sequence identity numbers (SEQ ID NOs.) for nucleotide and amino acid sequences referred to in the subject specification are defined after the bibliography. A summary of SEQ ID NOs. is also given in Table 1.

- 5 One aspect of the present invention contemplates an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding an amino acid sequence having homology to a regulator of gene expression or a derivative of said gene regulator.
- 10 Another aspect of the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a regulator of gene expression wherein said regulator comprises a zinc finger domain of an (HC<sub>3</sub>)<sub>2</sub> type.

Yet another aspect of the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

- (i) a nucleotide sequence set forth in SEQ ID NO:2;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:3;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
  - (iv) a nucleotide sequence capable of hybridizing under low stringency conditions at 42°C to the nucleotide sequence set forth in (i), (ii) or (iii).

The nucleotide sequence set forth in SEQ ID NO:2 defines the gene, mcg4. This gene encodes a product, MCG4, having an amino acid sequence set forth in SEQ ID NO:3.

Even yet another aspect of the present invention provides a genetic construct comprising a vector portion and an animal, more particularly a mammalian and even more particularly a human mcg4 gene portion, which mcg4 gene portion is capable of encoding an MCG4 polypeptide or a functional or immunologically interactive derivative thereof.

Still yet another aspect of the present invention contemplates a method of detecting a condition caused or facilitated by an aberration in mcg4, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg4 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.

Even still a further aspect of the present invention relates to a method of detecting a condition caused or facilitated by an aberration in mcg4, said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG4 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.

Another aspect of the present invention contemplates a method for detecting MCG4 or a derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG4 or its derivatives or homologues for a time and under conditions sufficient for an antibody-MCG4 complex to form, and then detecting said complex.

A further aspect of the present invention contemplates an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding an 20 amino acid sequence having homology to a guanine nucleotide exchange factor (GEF) or a derivative thereof.

Yet another aspect of the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

25

- (i) a nucleotide sequence set forth in SEQ ID NO:4 or 6;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:5 or 7;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
  - (iv) a nucleotide sequence capable of hybridizing under low stringency conditions to the

nucleotide sequence set forth in (i), (ii) or (iii).

The nucleotide sequence set forth in SEQ ID NO:4 or 6 defines the gene, mcg7. This gene encodes a product, MCG7, having an amino acid sequence set forth in SEQ ID NO:5 or 7.

5

Even yet another aspect of the present invention provides a genetic construct comprising a vector portion and an animal, more particularly a mammalian and even more particularly a human mcg7 gene portion, which mcg7 gene portion is capable of encoding an MCG7 polypeptide or a functional or immunologically interactive derivative thereof.

10

Still yet another aspect of the present invention contemplates a method of detecting a condition caused or facilitated by an aberration in mcg7, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg7 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.

Even still a further aspect of the present invention relates to a method of detecting a condition caused or facilitated by an aberration in mcg7, said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG7 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.

Another aspect of the present invention contemplates a method for detecting MCG7 or a derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG7 or its derivatives or homologues for a time and under conditions sufficient for an antibody-MCG7 complex to form, and then detecting said complex.

Yet another aspect of the present invention contemplates an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding an amino acid sequence having homology to a heat shock protein or a heat shock binding protein or a derivative thereof.

Another aspect of the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

- (i) a nucleotide sequence set forth in SEQ ID NO:8;
- 5 (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:9;
  - (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
  - (iv) a nucleotide sequence capable of hybridizing under low stringency conditions at 41°C to the nucleotide sequence set forth in (i), (ii) or (iii).

10

The nucleotide sequence set forth in SEQ ID NO:8 defines the gene, mcg18. This gene encodes a product, MCG18, having an amino acid sequence set forth in SEQ ID NO:7.

Even yet another aspect of the present invention provides a genetic construct comprising a vector portion and an animal, more particularly a mammalian and even more particularly a human mcg18 gene portion, which mcg18 gene portion is capable of encoding an MCG18 polypeptide or a functional or immunologically interactive derivative thereof.

Still yet another aspect of the present invention contemplates a method of detecting a condition caused or facilitated by an aberration in mcg18, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg18 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.

25

Even still a further aspect of the present invention relates to a method of detecting a condition caused or facilitated by an aberration in mcg18, said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG18 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.

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Another aspect of the present invention contemplates a method for detecting MCG18 or a

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derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG18 or its derivatives or homologues for a time and under conditions sufficient for an antibody-MCG18 complex to form, and then detecting said complex.

5

A summary of SEQ ID Nos. referred to in the subject specification is shown in Table 1.

TABLE 1
SUMMARY OF SEQ ID Nos.

5	SEQ ID NO.	DESCRIPTION		
	1	amino acid repeat sequence in DnaJ homologues		
	2	Nucleotide sequence of mcg4		
	3	amino acid sequence of MCG4		
	4	nucleotide sequence of mcg7		
10	5	amino acid sequence of MCG7		
	6	nucleotide sequence of mcg7 within exon of		
		nucleotides 183-288		
	7	amino acid sequence of MCG7 within exon of		
		nucleotide 183-288		
	8	nucleotide sequence of mcg18		
	9	amino acid sequence of MCG18		
15	10-18	amino acid sequence identified using BESTFIT		
	19	sequence of pGEX and mcg7 junction		
	20	sequence of pGEX and mcg7 junction		
	21	nucleotide sequence of myc-tag/mcg7 junction		
	22	amino acid sequence corresponding to SEQ ID NO:21		
20	23	nucleotide sequence of pGEX and mcg7 junction		
	24	amino acid sequence corresponding to SEQ ID NO:23		
	25-36	mcg7-specific oligonucleotide		
	37-45	mcg18-specific oligonucleotide		

<sup>25.</sup> Single and three letter abbreviations for amino acid residues are shown in Table 2.

TABLE 2

Amino Acid	Three-letter Abbreviation	One-letter Symbol
Alanine	Ala	A
Arginine	Arg	R
Asparagine	Asn	N
Aspartic acid	Asp	D
Cysteine	Cys	C
Glutamine	Gln	Q
Glutamic acid	Glu	E
Glycine	Gly	G
Histidine	His	Н
Isoleucine	Ile	I
Leucine	Leu	L
Lysine	Lys	K
Methionine	Met	M
Phenylalanine	Phe	F
Proline	Pro	P
Serine	Ser	S
Threonine	Thr	Т
Tryptophan	Trp	W
Tyrosine	Tyr	Y
Valine	Val	V
Any residue	Xaa	X

#### **BRIEF DESCRIPTION OF THE FIGURES**

Figure 1 is a representation of the nucleotide sequence [SEQ ID NO:2] and corresponding amino acid sequence [SEQ ID NO:3] of mcg4.

5

Figure 2 is a representation of the alignment of the human MCG4 amino acid sequence with a translation of a partial murine expressed sequence tag (EST).

Figure 3 is a representation of the alignment of the human MCG4 amino acid sequence with a 10 translation of a partial nematode EST.

Figure 4 is a diagrammatic representation showing a predicted structure of MCG4 where H and C represent histidine and cysteine residues, respectively and X refers to any amino acid residue. Zn represent zinc atoms.

15

Figure 5 is a representation of sensitive sequence homology search of related cysteine-containing motifs in another Caenorhabditis elegans protein.

Figure 6 is a representation showing that a related cysteine containing motif is present in the 20 GATA-binding transcription factor from Saccharomyces pombe.

Figure 7 is a Northern blot showing expression of mcg4 in various cultured human cancer cell lines. Lanes 1-5, respectively, represent the hybridization signal from 15µg total RNA derived from various human cancer cell lines. Lanes 1-5, respectively, contain RNA from H69 lung 25 carcinoma cells, JAM ovary carcinoma cells, BT20 breast carcinoma cells, HaCat transformed keratinocytes, T24 bladder carcinoma cells.

Figure 8 is a representation of a partial alignment of mcg4 with human ESTs AA074703 and AA134788.

30

Figure 9 is a representation of the partial nucleotide sequence alignment between a human

(W32939) and mouse (AA242159) mcg4-like EST in the putative 5' UTR of the mcg4 cDNA. The putative initiation codon is underlined and the region upstream represents 5' UTR.

Figure 10 is a representation showing MacVector alignment of MCG4 with forward translations
5 of ESTs AA134788 and AA074703. The nucleotide sequences are shown in Figure 8.

Figure 11 is a diagrammatic representation of the domains of MCG4

zinc finger consensus: CX<sub>2</sub>HX<sub>4</sub>CX<sub>2</sub>CX<sub>4</sub>HX<sub>2</sub>CX<sub>17</sub>CX<sub>2</sub>CX<sub>18</sub>HX<sub>2</sub>CX<sub>18</sub>CX<sub>2</sub>C

acidic domain consensus: 9/34 amino acids negatively charged, 0/34 positively charged

basic domain consensus: 13/55 amino acids positively charged, 0/55 negatively charged

leucine zipper domain consensus: LX<sub>6</sub>LX<sub>6</sub>RX<sub>6</sub>LX<sub>6</sub>L

alternate "novel" leucine zipper-like motif where leucine would not be aligned along the one surface of an alpha he!ix domain: (aa261) LX<sub>6</sub>LXLX<sub>6</sub>LXLX<sub>6</sub>L (aa 286).

15 Figure 12 is a representation showing similarity of MCG7 with GEFs of various organisms.

Figure 13(a) is a representation of the nucleotide sequence [SEQ ID NO:4] and corresponding amino acid sequence [SEQ ID NO:5] of mcg7. Nucleotides 183-288 are an alternative spliced exon (shown in lower case).

20

Figure 13(b) is a representation of the partial nucleotide sequence [SEQ ID NO:6] and corresponding amino acid sequence [SEQ ID NO:7] of mcg7 but without the exon shown in Fig. 13(a). Amino acids have been numbered from the first methionine codon (underlined). The cDNA molecules of Fig. 13(a) and Fig. 13(b) differ by the inclusion and exclusion of the exon 25 of nucleotides 183-288.

Figure 14 is a representation showing a comparison between MCG7 and a homologue from *Caenorhabditis elegans* using the BESTFIT algorithm. in the figure, the following sequences are underlined:

1a nematodeDVDEEDEVEDIEF [SEQ ID NO:10]1b humanDVDGDGHISQEEF [SEQ ID NO:11]nematodeDHDRDGFISQEEF [SEQ ID NO:12]

1c human DONODGCISREEM [SEQ ID NO:13]

5 nematode DVDMDGQISKDEL [SEQ ID NO:14]

#### GUANINE NT BINDING REGION = BLOCKS DATABASE NO. BL00720B

2 human HFVHVAEKLLQLQNFNTLMAVVGGLSHSSISRLKETH [SEQ ID NO:15]

nematode KFVHVAKHLRKINNFNTLMSVVGGITHSSVARLAKTY

10 [SEQ ID NO:16]

### DaG-PE BINDING DOMAIN = PROSITE DATABASE NO. PD0C00379

3 human HNFQESNSLRPVACRHCKALILGIYKQGLKCRACGVNCHKQCKDRLSVEC [SEQ ID NO:17]

15 nematode HNFHETTFLTPTTCNHCNKLLWGILRQGFKCKDCGLAVHSCCKSNAVAEC [SEQ ID NO:18]

Figure 15 is a representation of an alignment of human and a partial (5' UTR and partial coding sequence) murine mcg7 cDNA (GenBank Acc. No. W71787 and AA237373). The putative initiation codon is underlined. The murine sequence represents a composite of 2 partial cDNA sequences from the EST database (accession numbers W71787 and AA237373). Nucleotide differences between human and murine sequences are shown in lower case lettering and identical residues are indicated with asterisks.

- 25 Figure 16 is a representation of further 5' nucleotide and corresponding amino acid sequence for human mcg7. Nucleotide positions 1-321 were derived from GenBank Acc. No. AC000134 and nucleotides 322 onwards from Fig. 13(a). Two in-frame initiation codons are underlined. Asterisks denote in-frame stop codons.
- 30 Figure 17 is a graphical representation of a GDP release assay. □ Experiment #1 (mean of duplicates). ♦ Experiment #2 (mean of duplicates). The exchange reaction contained 36pmols

of GST-MCG (N-terminally truncated; encoded by Construct B in Fig. 18) and 1.6-12.8 pmols of recombinant GST-N-Ras.GDP. Reaction time 6 mins.

Estimated reaction constants:

 $K_m = 2.1 \mu M$ ,  $V_{max} = 37 p Mol/6 min/36 p Mol [Expt#1]$ 

5  $K_m = 1.5 \mu M$ ,  $V_{max} = 30.3 p Mol/6 min/36 p Mol [Expt#2]$ 

Figure 18 depicts various recombinant plasmids containing partial or full-length mcg7.

Figure 19 is a representation of the nucleotide sequence [SEQ ID NO:8] and corresponding amino acid sequence [SEQ ID NO:9] of mcg18.

Figure 20 is a representation showing that MCG18 has partial homology to E. coli DnaJ.

Figure 21 is a representation showing that MCG18 has homology to two *Caenorhabitis elegans* proteins.

Figure 22 is a representation showing that MCG18 has homology to a Saccharomyces pombe protein.

20 Figure 23 is a representation showing homology of MCG18 to a Drosophila virilis protein.

Figure 24 is a representation showing homology of MCG18 to human DnaJ proteins HDJ-2/HSDJ, HDJ-1/HSP40 and HSJ1.

25 Figure 25 is a representation of the nucleotide and corresponding amino acid sequence of murine mcg18.

Figure 26 is a representation of homology between human and murine MCG18.

30 Figure 27 depicts nucleotide sequences corresponding to the 5' untranslated region of human mcg18.

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Figure 28 depicts a Northern blot showing expression of mcg18 transcripts in total RNA isolated from various human cancer cell lines grown in culture. Lanes 1-5 respectively contain  $15\mu g$  RNA from H69 lung carcinoma cells, JAM ovary carcinoma cells, BT20 breast carcinoma cells, HaCat transformed keratinocytes, T24 bladder carcinoma cells.

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#### DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

The present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding an amino acid sequence having 5 homology to a regulator of gene expression or a derivative of said gene regulator.

More particularly, the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding a regulator of gene expression wherein said regulator comprises a zinc finger domain of an (HC<sub>3</sub>)<sub>2</sub> type.

Still more particularly, the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

- 15 (i) a nucleotide sequence set forth in SEQ ID NO:2;
  - (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:3;
  - (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridizing under low stringency conditions at 42°C to the nucleotide sequence set forth in (i), (ii) or (iii).

The present invention also provides an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding an amino acid sequence having homology to a guanine nucleotide exchange factor (GEF) or a derivative thereof.

More particularly, the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

- (i) a nucleotide sequence set forth in SEQ ID NO:4 or 6;
- a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:5 or 7;

- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridizing under low stringency conditions at 42°C to the nucleotide sequence set forth in (i), (ii) or (iii).

5

Another aspect of the present invention contemplates an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding an amino acid sequence having homology to a heat shock protein or a heat shock-binding protein or a derivative thereof.

10

More particularly, the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

- (i) a nucleotide sequence set forth in SEQ ID NO:8;
- 15 (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:9;
  - (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
  - (iv) a nucleotide sequence capable of hybridizing under low stringency conditions at 42°C to the nucleotide sequence set forth in (i), (ii) or (iii).

20

Preferably, the percentage similarity is at least about 50%. More preferably, the percentage similarity is at least about 60%.

Reference herein to a low stringency at 42°C includes and encompasses from at least about 1% v/v to at least about 15% v/v formamide and from at least about 1M to at least about 2M salt for hybridisation, and at least about 1M to at least about 2M salt for washing conditions. Alternative stringency conditions may be applied where necessary, such as medium stringency, which includes and encompasses from at least about 16% v/v to at least about 30% v/v formamide and from at least about 0.5M to at least about 0.9M salt for hybridisation, and at least about 0.5M and to at least about 0.9M salt for washing conditions, or high stringency, which includes and encompasses from at least about 31% v/v to at least about 50% v/v formamide and from at least

about 0.01M to at least about 0.15M salt for hybridisation, and at least about 0.01M to at least about 0.15M salt for washing conditions.

The term "similarity" as used herein includes exact identity between compared sequences at the 5 nucleotide or amino acid level. Where there is non-identity at the nucleotide level, "similarity" includes differences between sequences which result in different amino acids that are nevertheless related to each other at the structural, functional, biochemical and/or conformational levels. Where there is non-identity at the amino acid level, "similarity" includes amino acids that are nevertheless related to each other at the structural, functional, biochemical and/or conformational 10 levels.

The present invention extends to nucleic acid molecules with percentage similarities of approximately 65%, 70%, 75%, 80%, 85%, 90% or 95% or above or a percentage in between.

15 The nucleic acid molecule of the present invention defined by SEQ ID NO:2 is hereinafter referred to as constituting the "mcg4" gene. The protein encoded by mcg4 is referred to herein as "MCG4" and has an amino acid sequence set forth in SEQ ID NO:3. The mcg4 gene is proposed to encode, in accordance with the present invention, a regulator of gene expression and comprises a novel zinc finger domain, (HC<sub>3</sub>)<sub>2</sub>. A regulator of gene expression includes a 20 transcription factor. Regulation may be at the level of nucleic acid:protein or protein:protein interaction.

The nucleic acid molecule of the present invention defined by SEQ ID NO:4 or 6 is hereinafter referred to as constituting the "mcg7" gene. The protein encoded by mcg7 is referred to herein 25 as "MCG7" and has an amino acid sequence set forth in SEQ ID NO:5 or 7 and is involved in signal transduction. The difference in the nucleotide and amino acid sequence is due to the presence or absence of an exon at nucleotides 183-288.

The nucleic acid molecule of the present invention defined by SEQ ID NO:8 is hereinafter 30 referred to as constituting the "mcg18" gene. The protein encoded by mcg18 is referred to herein as "MCG18" and comprises the amino acid set forth in SEQ ID NO:9.

The present invention extends to the naturally occurring genomic mcg4, mcg7 and mcg18 nucleotide sequences or corresponding cDNA sequences or to derivatives thereof. Derivatives contemplated in the present invention include fragments, parts, portions, mutants, homologues and analogues of MCG4, MCG7 or MCG8 or the corresponding genetic sequences. Derivatives also include single or multiple amino acid substitutions, deletions and/or additions to MCG4, MCG7 or MCG18 or single or multiple nucleotide substitutions, deletions and/or additions to mcg4, mcg7 or mcg18. "Additions" to the amino acid or nucleotide sequences include fusions with other peptides, polypeptides or proteins or fusions to nucleotide sequences. Reference herein to "MCG4" or "mcg4", "MCG7" or "mcg7" or "MCG8" or mcg18" includes reference to all derivatives thereof including functional derivatives and immunologically interactive derivatives of MCG4, MCG7 or MCG18.

The mcg4, mcg7 and mcg18 of the present invention are particularly exemplified herein from humans and in particular from human chromosome 11q13.

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The present invention extends, however, to a range of homologues from, for example, primates, livestock animals (eg. sheep, cows, horses, donkeys, pigs), companion animals (eg. dogs, cats) laboratory test animals (eg. rabbits, mice, rats, guinea pigs), reptiles, birds (eg. chickens, ducks, geese, parrots), insects, nematodes, eukaryotic microorganisms and captive wild animals (eg. deer, foxes, kangaroos). Reference herein to mcg4 and mcg18 or their respective proteins MCG4, MCG7 and MCG18 includes reference to these molecules of human origin as well as novel forms of non-human origin.

The nucleic acid molecules of the present invention may be DNA or RNA. When the nucleic acid molecule is in DNA form, it may be genomic DNA or cDNA. RNA forms of the nucleic acid molecules of the present invention are generally mRNA.

Although the nucleic acid molecules of the present invention are generally in isolated form, they may be integrated into or ligated to or otherwise fused or associated with other genetic molecules such as vector molecules and in particular expression vector molecules. Vectors and expression vectors are generally capable of replication and, if applicable, expression in one or

both of a prokaryotic cell or a eukaryotic cell. Preferably, prokaryotic cells include *E. coli*, *Bacillus sp* and *Pseudomonas sp*. Preferred eukaryotic cells include yeast, fungal, mammalian and insect cells.

- 5 Accordingly, another aspect of the present invention contemplates a genetic construct comprising a vector portion and an animal, more particularly a mammalian and even more particularly a human mcg4 gene portion, which mcg4 gene portion is capable of encoding an MCG4 polypeptide or a functional or immunologically interactive derivative thereof.
- 10 Preferably, the *mcg4* gene portion of the genetic construct is operably linked to a promoter in the vector such that said promoter is capable of directing expression of said *mcg4* gene portion in an appropriate cell.

In addition, the *mcg4* gene portion of the genetic construct may comprise all or part of the gene 15 fused to another genetic sequence such as a nucleotide sequence encoding glutathione-Stransferase or part thereof.

The present invention extends to such genetic constructs and to prokaryotic or eukaryotic cells comprising same.

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It is proposed in accordance with the present invention that MCG4 is a transcription factor involved in gene regulation. Mutations in mcg4 may result in aberrations in gene regulation leading to the development of or a propensity to develop various types of cancer. In this regard, although not wishing to limit the present invention to any one hypothesis or mode of action, it 25 is proposed that mcg4 or its expression product may be involved in the tissue-specific or temporal regulation of particular genes.

A, deletion or aberration in the mcg4 gene may also be important in the detection of cancer or a propensity to develop cancer. An aberration may be a homozygous mutation or a heterozygous mutation. The detection may occur at the foetal or post-natal level. Detection may also be at the germline or somatic cell level. Furthermore, a risk of developing cancer may

be determined by assaying for aberrations in the parents and/or proband of a subject under investigation.

According to this aspect of the present invention, there is contemplated a method of detecting a condition caused or facilitated by an aberration in mcg4, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg4 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.

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Another aspect of the present invention contemplates a genetic construct comprising a vector portion and an animal, more particularly a mammalian and even more particularly a human mcg7 gene portion, which mcg7 gene portion is capable of encoding an mcg7 polypeptide or a functional or immunologically interactive derivative thereof.

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Preferably, the mcg7 gene portion of the genetic construct is operably linked to a promoter on the vector such that said promoter is capable of directing expression of said mcg7 gene portion in an appropriate cell.

20 In addition, the mcg7 gene portion of the genetic construct may comprise all or part of the gene fused to another genetic sequence such as a nucleotide sequence encoding glutathione-Stransferase or part thereof.

The present invention extends to such genetic constructs and to prokaryotic or eukaryotic cells comprising same.

It is proposed in accordance with the present invention that MCG7 is a GEF involved in signal transduction. Mutations in mcg7 or MCG7 may result in defective control of cell proliferation leading to the development of or a propensity to develop various types of cancer.

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A deletion or aberration in the mcg7 gene may also be important in the detection of cancer or

a propensity to develop cancer. An aberration may be a homozygous mutation or a heterozygous mutation. The detection may occur at the foetal or post-natal level. Detection may also be at the germline or somatic cell level. Furthermore, a risk of developing cancer may be determined by assaying for aberrations in the parents of a subject under investigation.

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According to this aspect of the present invention, there is contemplated a method of detecting a condition caused or facilitated by an aberration in mcg7, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg7 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.

Yet another aspect of the present invention contemplates a genetic construct comprising a vector portion and an animal, more particularly a mammalian and even more particularly a human 15 mcg18 gene portion, which mcg18 gene portion is capable of encoding an MCG18 polypeptide or a functional or immunologically interactive derivative thereof.

Preferably, the mcg18 gene portion of the genetic construct is operably linked to a promoter on the vector such that said promoter is capable of directing expression of said mcg18 gene portion 20 in an appropriate cell.

In addition, the *mcg18* gene portion of the genetic construct may comprise all or part of the gene fused to another genetic sequence such as a nucleotide sequence encoding glutathione-S-transferase or part thereof.

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The present invention extends to such genetic constructs and to prokaryotic or eukaryotic cells comprising same.

It is proposed in accordance with the present invention that MCG18 is a transcription factor 30 involved in protein folding, protein complex assembly and transit through subcellular compartments. MCG18 may also have a role in tumour suppression. Thus mutations in mcg18

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may result in the development of or a propensity to develop various types of cancer.

A deletion or aberration in the mcg18 gene may also be important in the detection of cancer or a propensity to develop cancer. An aberration may be a homozygous mutation or a 5 heterozygous mutation. The detection may occur at the foetal or post-natal level. Detection may also be at the germline or somatic cell level. Furthermore, a risk of developing cancer may be determined by assaying for aberrations in the parents and/or proband of the subject under investigation.

10 According to this aspect of the present invention, there is contemplated a method of detecting a condition caused or facilitated by an aberration in mcg18, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg18 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or 15 a propensity to develop said condition.

The nucleotide substitutions, additions or deletions may be detected by any convenient means including nucleotide sequencing, restriction fragment length polymorphism (RFLP), polymerase chain reaction (PCR), oligonucleotide hybridization and single stranded conformation polymorphism analysis (SSCP) amongst many others. An aberration includes modification to existing nucleotides such as to modify glycosylation signal amongst other effects.

In an alternative method, aberrations in the mcg4, mcg7 and mcg18 genes are detected by screening for mutations in MCG4, MCG7 and MCG18, respectively.

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A mutation in MCG4, MCG7 or MCG18 may be a single or multiple amino acid substitution, addition and/or deletion. The mutation in mcg4, mcg7 or mcg18 may also result in either no translation product being produced or a product in truncated form. A mutant may also be an altered glycosylation pattern or the introduction of side chain modifications to amino acid residues.

According to this aspect of the present invention, there is provided a method of detecting a condition caused or facilitated by an aberration in mcg4, mcg7 or mcg18 said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG4, MCG7 or MCG18 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.

A particularly convenient means of detecting a mutation in MCG4, MCG7 or MCG18 is by use of antibodies.

- 10 Accordingly another aspect of the present invention is directed to antibodies to MCG4, MCG7 or MCG18 and its derivatives. Such antibodies may be monoclonal or polyclonal and may be selected from naturally occurring antibodies to MCG4, MCG7 or MCG18 or may be specifically raised to MCG4, MCG7 or MCG18 or derivatives thereof. In the case of the latter, MCG4, MCG7 or MCG18 or their derivatives may first need to be associated with a carrier molecule.
- 15 The antibodies to MCG4, MCG7 or MCG18 of the present invention are particularly useful as diagnostic agents.

For example, antibodies to MCG4, MCG7 or MCG18 and their derivatives can be used to screen for wild-type MCG4, MCG7 or MCG18 or for mutated MCG4, MCG7 or MCG18 molecules.

- 20 The latter may occur, for example, during or prior to certain cancer development. A differential binding assay is also particularly useful. Techniques for such assays are well known in the art and include, for example, sandwich assays and ELISA. Knowledge of normal MCG4, MCG7 or MCG18 levels or the presence of wild-type MCG4, MCG7 or MCG18 may be important for diagnosis of certain cancers or a predisposition for development of cancers or for monitoring 25 certain therapeutic protocols.
- As stated above antibodies to MCG4, MCG7 or MCG18 of the present invention may be monoclonal or polyclonal or may be fragments of antibodies such as Fab fragments. Furthermore, the present invention extends to recombinant and synthetic antibodies and to antibody hybrids. A "synthetic antibody" is considered herein to include fragments and hybrids of antibodies.

For example, specific antibodies can be used to screen for wild-type MCG4, MCG7 or MCG18 molecule or specific mutant molecules such as molecules having a certain deletion. This would be important, for example, as a means for screening for levels of MCG4, MCG7 or MCG18 in a cell extract or other biological fluid or purifying MCG4, MCG7 or MCG18 made by recombinant means from culture supernatant fluid or purified from a cell extract. Techniques for the assays contemplated herein are known in the art and include, for example, sandwich assays and ELISA.

It is within the scope of this invention to include any second antibodies (monoclonal, polyclonal or fragments of antibodies or synthetic antibodies) directed to the first mentioned antibodies discussed above. Both the first and second antibodies may be used in detection assays or a first antibody may be used with a commercially available anti-immunoglobulin antibody. An antibody as contemplated herein includes any antibody specific to any region of wild-type MCG4, MCG7 or MCG18 or to a specific mutant phenotype or to a deleted or otherwise altered region.

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Both polyclonal and monoclonal antibodies are obtainable by immunization of a suitable animal or bird with MCG4, MCG7 or MCG18 or its derivatives and either type is utilizable for immunoassays. The methods of obtaining both types of sera are well known in the art. Polyclonal sera are less preferred but are relatively easily prepared by injection of a suitable laboratory animal or bird with an effective amount of MCG4, MCG7 or MCG18 or antigenic parts thereof or derivatives thereof, collecting serum from the animal or bird, and isolating specific sera by any of the known immunoadsorbent techniques. Although antibodies produced by this method are utilizable in virtually any type of immunoassay, they are generally less favoured because of the potential heterogeneity of the product.

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The use of monoclonal antibodies in an immunoassay is particularly preferred because of the ability to produce them in large quantities and the homogeneity of the product. The preparation of hybridoma cell lines for monoclonal antibody production derived by fusing an immortal cell line and lymphocytes sensitized against the immunogenic preparation can be done by techniques which are well known to those who are skilled in the art.

Another aspect of the present invention contemplates a method for detecting MCG4, MCG7 or MCG18 or a derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG4, MCG7 or MCG18 or its derivatives or homologues for a time and under conditions sufficient for an antibody-MCG4, MCG7 or MCG18 complex to form, and then detecting said complex.

Preferably, the biological sample is a cell extract from a human or other animal or a bird.

The presence of MCG4, MCG7 or MCG18 may be accomplished in a number of ways such as by Western blotting and ELISA procedures. A wide range of immunoassay techniques are available as can be seen by reference to US Patent Nos. 4,016,043, 4, 424,279 and 4,018,653. These include both single-site and two-site or "sandwich" assays of the non-competitive types, as well as traditional competitive binding assays. These assays also include direct binding of a labelled antibody to a target.

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Sandwich assays are among the most useful and commonly used assays and are favoured for use in the present invention. A number of variations of the sandwich assay technique exist, and all are intended to be encompassed by the present invention. Briefly, in a typical forward assay, an unlabelled antibody is immobilized on a solid substrate and the sample to be tested brought into 20 contact with the bound molecule. After a suitable period of incubation, for a period of time sufficient to allow formation of an antibody-antigen complex, a second antibody specific to the antigen, labelled with a reporter molecule capable of producing a detectable signal is then added and incubated, allowing time sufficient for the formation of another complex of antibody-antigenlabelled antibody. Any unreacted material is washed away, and the presence of the antigen is 25 determined by observation of a signal produced by the reporter molecule. The results may either be qualitative, by simple observation of the visible signal, or may be quantitated by comparing with a control sample containing known amounts of hapten. Variations on the forward assay include a simultaneous assay, in which both samrle and labelled antibody are added simultaneously to the bound antibody. These techniques are well known to those skilled in the 30 art, including any minor variations as will be readily apparent. In accordance with the present invention the sample is one which might contain MCG4, MCG7 or MCG18 including cell extract or tissue biopsy. The sample is, therefore, generally a biological sample comprising biological fluid but also extends to fermentation fluid and supernatant fluid such as from a cell culture.

In the typical forward sandwich assay, a first antibody having specificity for the MCG4, MCG7

or MCG18 or an antigenic part thereof or a derivative thereof or antigenic parts thereof, is either covalently or passively bound to a solid surface. The solid surface is typically glass or a polymer, the most commonly used polymers being cellulose, polyacrylamide, nylon, polystyrene, polyvinyl chloride or polypropylene. The solid supports may be in the form of tubes, beads, discs of microplates, or any other surface suitable for conducting an immunoassay. The binding processes are well-known in the art and generally consist of cross-linking covalently binding or physically adsorbing, the polymer-antibody complex is washed in preparation for the test sample. An aliquot of the sample to be tested is then added to the solid phase complex and incubated for a period of time sufficient (e.g. 2-40 minutes or overnight if more convenient) and under suitable conditions (e.g. from room temperature to 37°C) to allow binding of any subunit present in the antibody. Following the incubation period, the antibody subunit solid phase is washed and dried and incubated with a second antibody specific for a portion of the hapten. The second antibody is linked to a reporter molecule which is used to indicate the binding of the second antibody to the hapten.

20 An alternative method involves immobilizing the target molecules in the biological sample and then exposing the immobilized target to specific antibody which may or may not be labelled with a reporter molecule. Depending on the amount of target and the strength of the reporter molecule signal, a bound target may be detectable by direct labelling with the antibody. Alternatively, a second labelled antibody, specific to the first antibody is exposed to the target-first antibody complex to form a target-first antibody-second antibody tertiary complex. The complex is detected by the signal emitted by the reporter molecule.

By "reporter molecule" as used in the present specification, is meant a molecule which, by its chemical nature, provides an analytically identifiable signal which allows the detection of antigen-bound antibody. Detection may be either qualitative or quantitative. The most commonly used reporter molecules in this type of assay are either enzymes, fluorophores or radionuclide

containing molecules (i.e. radioisotopes) and chemiluminescent molecules.

In the case of an enzyme immunoassay, an enzyme is conjugated to the second antibody, generally by means of glutaraldehyde or periodate. As will be readily recognized, however, a wide variety of different conjugation techniques exist, which are readily available to the skilled 5 artisan. Commonly used enzymes include horseradish peroxidase, glucose oxidase, betagalactosidase and alkaline phosphatase, amongst others. The substrates to be used with the specific enzymes are generally chosen for the production, upon hydrolysis by the corresponding enzyme, of a detectable colour change. Examples of suitable enzymes include alkaline phosphatase and peroxidase. It is also possible to employ fluorogenic substrates, which yield a 10 fluorescent product rather than the chromogenic substrates noted above. In all cases, the enzyme-labelled antibody is added to the first antibody hapten complex, allowed to bind, and then the excess reagent is washed away. A solution containing the appropriate substrate is then added to the complex of antibody-antigen-antibody. The substrate will react with the enzyme linked to the second antibody, giving a qualitative visual signal, which may be further quantitated, 15 usually spectrophotometrically, to give an indication of the amount of hapten which was present in the sample. "Reporter molecule" also extends to use of cell agglutination or inhibition of agglutination such as red blood cells on latex beads, and the like.

Alternately, fluorescent compounds, such as fluorescein and rhodamine, may be chemically coupled to antibodies without altering their binding capacity. When activated by illumination with light of a particular wavelength, the fluorochrome-labelled antibody adsorbs the light energy, inducing a state to excitability in the molecule, followed by emission of the light at a characteristic colour visually detectable with a light microscope. As in the EIA, the fluorescent labelled antibody is allowed to bind to the first antibody-hapten complex. After washing off the unbound reagent, the remaining tertiary complex is then exposed to the light of the appropriate wavelength the fluorescence observed indicates the presence of the hapten of interest. Immunofluorescence and EIA techniques are both very well established in the art and are particularly preferred for the present method. However, other reporter molecules, such as radioisotope, chemiluminescent or bioluminescent molecules, may also be employed.

MCG7 or MCG18 or functional derivatives thereof. Such genetic constructs are also contemplated to be useful in modulating expression of specific genes in which mcg4, mcg7 or mcg18 is involved in tissue-specific or temporal regulation.

- 5 Accordingly, another aspect of the present invention is directed to a genetic construct comprising a nucleotide sequence encoding a peptide, polypeptide or protein and mcg4, mcg7 or mcg18 or a functional derivative or homologue thereof capable of modulating the expression of said nucleotide sequence.
- 10 As stated above, MCG18 is proposed to have a role in tumour suppression. Accordingly, it is further proposed in accordance with the present invention to use recombinant MCG18 in pharmaceutical preparations for treating arresting or otherwise ameliorating the effects of certain cancers.
- 15 Accordingly, another aspect of the present invention contemplates a method for treating, arresting or otherwise ameliorating the effects of a cancer in an animal or bird, said method comprising administering to said animal or bird an effective amount of MCG18 or a functional derivative thereof for a time and under conditions sufficient to treat, arrest or otherwise ameliorate the effects of said cancer.

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The present invention, therefore, contemplates a pharmaceutical composition comprising MCG18 or a derivative thereof or a modulator of mcg18 expression or MCG18 activity and one or more pharmaceutically acceptable carriers and/or diluents. These components are referred to hereinafter as the "active ingredients". The active ingredients may also include anti-cancer agents or agents which facilitate actions of MCG18.

The pharmaceutical forms suitable for injectable use include sterile aqueous solutions (where water soluble) and sterile powders for the extemporaneous preparation of sterile injectable solutions. It must be stable under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms such as bacteria and fungi. The carrier may be a solvent medium containing, for example, water, ethanol, polyol (for example,

glycerol, propylene glycol and liquid polyethylene glycol, and the like), suitable mixtures thereof, and vegetable oils. The proper fluidity can be maintained, for example, by the use of a coating such as licithin and by the use of superfactants. The preventions of the action of microorganisms can be brought about by various antibacterial and antifungal agents, for example, parabens, 5 chlorobutanol, phenol, sorbic acid, thimersal and the like. In many cases, it will be preferable to include isotonic agents, for example, sugars or sodium chloride. Prolonged absorption of the injectable compositions can be brought about by the use in the compositions of agents delaying absorption, for example, aluminum monostearate and gelatin.

10 Sterile injectable solutions are prepared by incorporating the active compounds in the required amount in the appropriate solvent with various of the other ingredients enumerated above, as required, followed by filtered sterilization. In the case of sterile powders for the preparation of sterile injectable solutions, the preferred methods of preparation are vacuum drying and the freeze-drying technique which yield a powder of the active ingredient plus any additional desired ingredient from previously sterile-filtered solution thereof.

When the active ingredients are suitably protected they may be orally administered, for example, with an inert diluent or with an assimilable edible carrier, or it may be enclosed in hard or soft shell gelatin capsule, or it may be compressed into tablets, or it may be incorporated directly with the food of the diet. For oral therapeutic administration, the active compound may be incorporated with excipients and used in the form of ingestible tablets, buccal tablets, troches, capsules, elixirs, suspensions, syrups, wafers, and the like. Such compositions and preparations should contain at least 1% by weight of active compound. The percentage of the compositions and preparations may, of course, be varied and may conveniently be between about 5 to about 80% of the weight of the unit. The amount of active compound in such therapeutically useful compositions in such that a suitable dosage will be obtained. Preferred compositions or preparations according to the present invention are prepared so that an oral dosage unit form contains between about 0.1 µg and 2000 mg of active compound.

30 The tablets, troches, pills, capsules and the like may also contain the components as listed hereafter. A binder such as gum, acacia, corn starch or gelatin; excipients such as dicalcium

phosphate; a disintegrating agent such as corn starch, potato starch, alginic acid and the like; a lubricant such as magnesium stearate; and a sweetening agent such a sucrose, lactose or saccharin may be added or a flavouring agent such as peppermint, oil of wintergreen, or cherry flavouring. When the dosage unit form is a capsule, it may contain, in addition to materials of the above type, a liquid carrier. Various other materials may be present as coatings or to otherwise modify the physical form of the dosage unit. For instance, tablets, pills, or capsules may be coated with shellac, sugar or both. A syrup or elixir may contain the active compound, sucrose as a sweetening agent, methyl and propylparabens as preservatives, a dye and flavouring such as cherry or orange flavour. Of course, any material used in preparing any dosage unit form should be pharmaceutically pure and substantially non-toxic in the amounts employed. In addition, the active compound(s) may be incorporated into sustained-release preparations and formulations.

The present invention also extends to forms suitable for topical application such as creams, lotions and gels.

Pharmaceutically acceptable carriers and/or diluents include any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents and the like. The use of such media and agents for pharmaceutical active substances is well known in the art. Except insofar as any conventional media or agent is incompatible with the active ingredient, use thereof in the therapeutic compositions is contemplated. Supplementary active ingredients can also be incorporated into the compositions.

It is especially advantageous to formulate parenteral compositions in dosage unit form for ease of administration and uniformity of dosage. Dosage unit form as used herein refers to physically discrete units suited as unitary dosages for the mammalian subjects to be treated; each unit containing a predetermined quantity of active material calculated to produce the desired theraperatic effect in association with the required pharmaceutical carrier. The specification for the novel dosage unit forms of the invention are dictated by and directly dependent on (a) the unique characteristics of the active material and the particular therapeutic effect to be achieved, and (b) the limitations inherent in the art of compounding such an active material for the

treatment of disease in living subjects having a diseased condition in which bodily health is impaired as herein disclosed in detail.

The principal active ingredient is compounded for convenient and effective administration in effective amounts with a suitable pharmaceutically acceptable carrier in dosage unit form as hereinbefore disclosed. A unit dosage form can, for example, contain the principal active compound in amounts ranging from 0.5 µg to about 2000 mg. Expressed in proportions, the active compound is generally present in from about 0.5 µg to about 2000 mg/ml of carrier. In the case of compositions containing supplementary active ingredients, the dosages are determined by reference to the usual dose and manner of administration of the said ingredients.

Effective amounts contemplated by the present invention include those amounts effective to ameliorate a condition. For example, it is envisaged that effective amounts would range from about  $0.001~\mu g/kg$  body weight to about 100~mg/kg body weight. Alternatively, effective 15 amounts of about  $0.01~\mu g/kg$  body weight to about 10~mg/kg body weight or even  $0.1~\mu g/kg$  body weight to about 1~mg/kg body weight. Administration may be per minute, hour, day, week, month or year or may only be a once off administration.

The pharmaceutical composition may also comprise genetic molecules such as a vector capable of transfecting target cells where the vector carries a nucleic acid molecule capable of modulating mcg18 expression or MCG18 activity. The vector may, for example, be a viral vector.

As stated above, the present invention further contemplates a range of derivatives of MCG18. Derivatives include fragments, parts, portions, mutants, homologues and analogues of the MCG18 polypeptide and corresponding genetic sequence. Derivatives also include single or multiple amino acid substitutions, deletions and/or additions to MCG18 or single or multiple nucleotide substitutions, deletions and/or additions to the genetic sequence encoding MCG18. "Additions" to amino acid sequences or nucleotide sequences include fusions with other peptides, polypeptides or proteins or fusions to nucleotide sequences. Reference herein to "MCG18" includes reference to all derivatives thereof including functional derivatives or MCG18 immunologically interactive derivatives.

Analogues of MCG18 contemplated herein include, but are not limited to, modification to side chains, incorporating of unnatural amino acids and/or their derivatives during peptide, polypeptide or protein synthesis and the use of crosslinkers and other methods which impose conformational constraints on the proteinaceous molecule or their analogues.

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Examples of side chain modifications contemplated by the present invention include modifications of amino groups such as by reductive alkylation by reaction with an aldehyde followed by reduction with NaBH<sub>4</sub>; amidination with methylacetimidate; acylation with acetic anhydride; carbamoylation of amino groups with cyanate; trinitrobenzylation of amino groups with 2, 4, 6-trinitrobenzene sulphonic acid (TNBS); acylation of amino groups with succinic anhydride and tetrahydrophthalic anhydride; and pyridoxylation of lysine with pyridoxal-5-phosphate followed by reduction with NaBH<sub>4</sub>.

The guanidine group of arginine residues may be modified by the formation of heterocyclic condensation products with reagents such as 2,3-butanedione, phenylglyoxal and glyoxal.

The carboxyl group may be modified by carbodiimide activation *via* O-acylisourea formation followed by subsequent derivitisation, for example, to a corresponding amide.

20 Sulphydryl groups may be modified by methods such as carboxymethylation with iodoacetic acid or iodoacetamide; performic acid oxidation to cysteic acid; formation of a mixed disulphides with other thiol compounds; reaction with maleimide, maleic anhydride or other substituted maleimide; formation of mercurial derivatives using 4-chloromercuribenzoate, 4-chloromercuriphenylsulphonic acid, phenylmercury chloride, 2-chloromercuri-4-nitrophenol and other mercurials; carbamoylation with cyanate at alkaline pH.

Tryptophan residues may be modified by, for example, oxidation with N-bromosuccinimide or alkylation of the indole ring with 2-hydroxy-5-nitrobenzyl bromide co sulphenyl halides.

Tyrosine residues on the other hand, may be altered by nitration with tetranitromethane to form 30 a 3-nitrotyrosine derivative.

Modification of the imidazole ring of a histidine residue may be accomplished by alkylation with iodoacetic acid derivatives or N-carbethoxylation with diethylpyrocarbonate.

Examples of incorporating unnatural amino acids and derivatives during peptide synthesis include, but are not limited to, use of norleucine, 4-amino butyric acid, 4-amino-3-hydroxy-5-phenylpentanoic acid, 6-aminohexanoic acid, t-butylglycine, norvaline, phenylglycine, ornithine, sarcosine, 4-amino-3-hydroxy-6-methylheptanoic acid, 2-thienyl alanine and/or D-isomers of amino acids. A list of unnatural amino acids, contemplated herein is shown in Table 3.

TABLE 3

5	Non-conventional amino acid	Code	Non-conventional amino acid	Code
	α-aminobutyric acid	Abu	L-N-methylalanine	Nmala
	α-amino-α-methylbutyrate	Mgabu	L-N-methylarginine	Nmarg
	aminocyclopropane-	Cpro	L-N-methylasparagine	Nmasn
	carboxylate		L-N-methylaspartic acid	Nmasp
)	aminoisobutyric acid	Aib	L-N-methylcysteine	Nmcys
	aminonorbornyl-	Norb	L-N-methylglutamine	Nmgln
	carboxylate		L-N-methylglutamic acid	Nmglu
	cyclohexylalanine	Chexa	L-N-methylhistidine	Nmhis
	cyclopentylalanine	Cpen	L-N-methylisolleucine	Nmile
5	D-alanine	Dal	L-N-methylleucine	Nmleu
	D-arginine	Darg	L-N-methyllysine	Nmlys
	D-aspartic acid	Dasp	L-N-methylmethionine	Nmmet
	D-cysteine	Dcys	L-N-methylnorleucine	Nmnle
	D-glutamine	Dgln	L-N-methylnorvaline	Nmnva
0	D-glutamic acid	Dglu	L-N-methylornithine	Nmorn
	D-histidine	Dhis	L-N-methylphenylalanine	Nmphe
	D-isoleucine	Dile	L-N-methylproline	Nmpro
	D-leucine	Dleu	L-N-methylserine	Nmser
	D-lysine	Dlys	L-N-methylthreonine	Nmthr
5	D-methionine	Dmet	L-N-methyltryptophan	Nmtrp
	D-ornithine	Dorn	L-N-methyltyrosine	Nmtyr
	D-phenylalanine	Dphe	L-N-methylvaline	Nmval
	D-proline	Dpro	L-N-methylethylglycine	Nmetg
1	D-serine	Dser	L-N-methyl-t-butylglycine	Nmtbug
0	D-threonine	Dthr	L-norleucine	Nle
	D-tryptophan	Dtrp	L-norvaline	Nva

	D-tyrosine	Dtyr	α-methyl-aminoisobutyrate	Maib
	D-valine	Dval	$\alpha$ -methyl- $\gamma$ -aminobutyrate	Mgabu
	D-α-methylalanine	Dmala	α-methylcyclohexylalanine	Mchexa
	D-α-methylarginine	Dmarg	α-methylcylcopentylalanine	Mcpen
5	D-α-methylasparagine	Dmasn	$\alpha$ -methyl- $\alpha$ -napthylalanine	Manap
	D-α-methylaspartate	Dmasp	α-methylpenicillamine	Mpen
	D-α-methylcysteine	Dmcys	N-(4-aminobutyl)glycine	Nglu
	$D$ - $\alpha$ -methylglutamine	Dmgln	N-(2-aminoethyl)glycine	Naeg
	$D$ - $\alpha$ -methylhistidine	Dmhis	N-(3-aminopropyl)glycine	Norn
10	D-α-methylisoleucine	Dmile	N-amino-α-methylbutyrate	Nmaabu
	D-α-methylleucine	Dmleu	α-napthylalanine	Anap
	D-α-methyllysine	Dmlys	N-benzylglycine	Nphe
	$D$ - $\alpha$ -methylmethionine	Dmmet	N-(2-carbamylethyl)glycine	Ngln
	$D$ - $\alpha$ -methylornithine	Dmorn	N-(carbamylmethyl)glycine	Nasn
15	$D$ - $\alpha$ -methylphenylalanine	Dmphe	N-(2-carboxyethyl)glycine	Nglu
	D-α-methylproline	Dmpro	N-(carboxymethyl)glycine	Nasp
	D-α-methylserine	Dmser	N-cyclobutylglycine	Ncbut
	D-α-methylthreonine	Dmthr	N-cycloheptylglycine	Nchep
	$D$ - $\alpha$ -methyltryptophan	Dmtrp	N-cyclohexylglycine	Nchex
20	D-α-methyltyrosine	Dmty	N-cyclodecylglycine	Ncdec
	D-α-methylvaline	Dmval	N-cylcododecylglycine	Ncdod
	D-N-methylalanine	Dnmala	N-cyclooctylglycine	Ncoct
	D-N-methylarginine	Dnmarg	N-cyclopropylglycine	Ncpro
	D-N-methylasparagine	Dnmasn	N-cycloundecylglycine	Ncund
25	D-N-methylaspartate	Dnmasp	N-(2,2-diphenylethyl)glycine	Nbhm
	D-N-methylcysteine	Dnmcys	N-(3,3-diphenylpropyl)glycine	Nbhe
	D-N-methylglutamine	Dnmgln	N-(3-guanidinopropyl)glycine	Narg
	D-N-methylglutamate	Dnmglu	N-(1-hydroxyethyl)glycine	Nthr
	D-N-methylhistidine	Dnmhis	N-(hydroxyethyl))glycine	Nser
30	D-N-methylisoleucine	Dnmile	N-(imidazolylethyl))glycine	Nhis
	D-N-methylleucine	Dnmleu	N-(3-indolylyethyl)glycine	Nhtrp

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	D-N-methyllysine	Dnmlys	N-methyl-γ-aminobutyrate	Nmgabu
	N-methylcyclohexylalanine	Nmchexa	D-N-methylmethionine	Dommet
	D-N-methylornithine	Dnmorn	N-methylcyclopentylalanine	Nmcpen
	N-methylglycine	Nala	D-N-methylphenylalanine	Dnmphe
5	N-methylaminoisobutyrate	Nmaib	D-N-methylproline	Dnmpro
	N-(1-methylpropyl)glycine	Nile	D-N-methylserine	Dnmser
	N-(2-methylpropyl)glycine	Nleu	D-N-methylthreonine	Dnmthr
	D-N-methyltryptophan	Dnmtrp	N-(1-methylethyl)glycine	Nval
	D-N-methyltyrosine	Dnmtyr	N-methyla-napthylalanine	Nmanap
10	D-N-methylvaline	Dnmval	N-methylpenicillamine	Nmpen
	γ-aminobutyric acid	Gabu	N-(p-hydroxyphenyl)glycine	Nhtyr
	L-t-butylglycine	Tbug	N-(thiomethyl)glycine	Ncys
	L-ethylglycine	Etg	penicillamine	Pen
	L-homophenylalanine	Hphe	L-α-methylalanine	Mala
15	$L$ - $\alpha$ -methylarginine	Marg	L-α-methylasparagine	Masn
	L-α-methylaspartate	Masp	L-α-methyl-t-butylglycine	Mtbug
	L-α-methylcysteine	Mcys	L-methylethylglycine	Metg
	$L$ - $\alpha$ -methylglutamine	Mgln	L-α-methylglutamate	Mglu
	L-α-methylhistidine	Mhis	L-α-methylhomophenylalanine	Mhphe
20	L-α-methylisoleucine	Mile	N-(2-methylthioethyl)glycine	Nmet
	L-α-methylleucine	Mleu	L-α-methyllysine	Mlys
	$L$ - $\alpha$ -methylmethionine	Mmet	L-α-methylnorleucine	Mnle
	L-α-methylnorvaline	Mnva	L-α-methylornithine	Morn
	L-α-methylphenylalanine	Mphe	L-α-methylproline	Mpro
25	L-α-methylserine	Mser	L-α-methylthreonine	Mthr
	L-α-methyltryptophan	Mtrp	L-α-methyltyrosine	Mtyr

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L-α-methylvaline

Mval

L-N-methylhomophenylalanine

Nmhphe

N-(N-(2,2-diphenylethyl)

Nnbhm

N-(N-(3,3-diphenylpropyl)

Nnbhe

carbamylmethyl)glycine

carbamylmethyl)glycine

1-carboxy-1-(2,2-diphenyl-

Nmbc

5 ethylamino)cyclopropane

Crosslinkers can be used, for example, to stabilise 3D conformations, using homo-bifunctional crosslinkers such as the bifunctional imido esters having (CH<sub>2</sub>)<sub>n</sub> spacer groups with n=1 to n=6, 10 glutaraldehyde, N-hydroxysuccinimide esters and hetero-bifunctional reagents which usually contain an amino-reactive moiety such as N-hydroxysuccinimide and another group specificreactive moiety such as maleimido or dithio moiety (SH) or carbodiimide (COOH). In addition, peptides can be conformationally constrained by, for example, incorporation of  $C_{\alpha}$  and  $N_{\alpha}$  methylamino acids, introduction of double bonds between  $C_{\alpha}$  and  $C_{\beta}$  atoms of amino acids and 15 the formation of cyclic peptides or analogues by introducing covalent bonds such as forming an amide bond between the N and C termini, between two side chains or between a side chain and the N or C terminus.

Such analogues also apply in respect of MCG4 and MCG7.

20

The present invention further contemplates chemical analogues of MCG18 capable of acting as antagonists or agonists of MCG18 or which can act as functional analogues of MCG18. Chemical analogues may not necessarily be derived from MCG18 but may share certain conformational similarities. Alternatively, chemical analogues may be specifically designed to 25 mimic certain physiochemical properties of MCG18. Chemical analogues may be chemically synthesised or may be detected following, for example, natural product screening.

The identification of MCG:8 permits the generation of a range of therapeutic molecules capable of modulating expression of MCG18 or modulating the activity of MCG18. Modulators 30 contemplated by the present invention includes agonists and antagonists of MCG18 expression. Antagonists of MCG18 expression include antisense molecules, ribozymes and co-suppression molecules. Agonists include molecules which increase promoter ability or interfere with negative regulatory mechanisms. Agonists of MCG18 include molecules which overcome any negative regulatory mechanism. Antagonists of MCG18 include antibodies and inhibitor peptide fragments.

5

These types of modifications may be important to stabilise MCG18 if administered to an individual or for use as a diagnostic reagent.

Other derivatives contemplated by the present invention include a range of glycosylation variants 10 from a completely unglycosylated molecule to a modified glycosylated molecule. Altered glycosylation patterns may result from expression of recombinant molecules in different host cells.

Another embodiment of the present invention contemplates a method for modulating expression of MCG18 in a human, said method comprising contacting the mcg18 gene encoding MCG18 with an effective amount of a modulator of mcg18 expression for a time and under conditions sufficient to up-regulate or down-regulate or otherwise modulate expression of mcg18. For example, a nucleic acid molecule encoding MCG18 or a derivative thereof may be introduced into a cell to facilitate protection of that cell from becoming cancerous.

20

Another aspect of the present invention contemplates a method of modulating activity of MCG18 in a human, said method comprising administering to said mammal a modulating effective amount of a molecule for a time and under conditions sufficient to increase or decrease MCG18 activity. The molecule may be a proteinaceous molecule or a chemical entity and may also be a derivative of MCG18 or a chemical analogue or truncation mutant of MCG18.

The present invention is further described with reference to the following non-limiting Examples.

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#### **EXAMPLE 1**

A human gene (designated mcg4) was identified on chromosome 11q13 that on the basis of sequence homology is predicted to encode a putative transcription factor of 310 amino acids 5 (Fig. 1). mcg4 is transcribed in several different cell lines (Fig. 7).

#### **EXAMPLE 2**

The expressed sequence tag (EST) database contains partial sequence data for the murine (Fig. 10 2) and nematode (Fig. 3) homologues of mcg4.

#### **EXAMPLE 3**

MCG4 contains a sequence of cysteine residues within the N-terminal region of the protein that resembles zinc-finger binding domains of a novel type, ie. (HC<sub>3</sub>)<sub>2</sub> [Fig. 4].

#### **EXAMPLE 4**

Sensitive sequence homology searches reveal that related cysteine-containing motifs are present 20 in another C. elegans protein (Fig. 5) as well as the GATA-binding transcription factor from S. pombe (Fig. 6).

#### **EXAMPLE 5**

25 mcg4 will have commercial value due to its likelihood of encoding a novel transcription factor that is highly conserved amongst organisms, thus suggesting an integral role in gene regulation. mcg4 may also be involved in some way in tissue-specific or temporal regulation of certain genes, thus making it a potential target for modulating expression of those downstream effectors.

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## **EXAMPLE 6**

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Nucleotide sequence data generated from cosmid clone cSRL-72c4 with the T7 primer (Promega, and Applied Biosystems Incorporated dye terminator sequencing kit) was aligned to the GenBank Expressed Sequence Tag (EST) database using the program BLASTN (Altschul et al 1990) and was found to match numerous human and mouse entries (Table 4 and Figure 2). These matching ESTs were further used to identify overlapping entries in the EST database (Table 5). The nucleotide sequences of these human ESTs were complied using MacVector 4.2.1 software (IBI-Kodak) to produce the cDNA sequence shown in Figure 1. EST entries AA074703 and AA134788 are closely related at the nucleotide level to mcg4 and it is, therefore, likely that mcg4 is a member of a newly discovered gene family (Figure 8).

The cDNA sequence of mcg4 was translated in all possible reading frames and compared to the GenBank non-redundant protein database using the program BLASTX (Altschul et al, 1990) at 15 the National Center for Biotechnology Information (http://www.ncbi.nih.gov.nlm). As the protein appeared to be novel, a translation of the longest reading frame for the mcg4 cDNA was aligned to the EST database using the program TBLASTN, which performed a dynamic translation of the EST database in all 6 frames. The search results indicated that the nematode C. elegans had an MCG4-like protein (Figure 3), with the matching domains containing a spatial 20 sequence of Cysteine and Histidine residues which resembled a zinc-finger structure (Figure 4). The program BLASTP was used, therefore, to conduct sensitive searches of the protein databases for similar zinc-finger motifs. A weak match to the putative zinc-finger domain was observed for another protein from C. elegans (Figure 5) and a poorer match for the GATAbinding transcription factor from S. pombe (Figure 6). The putative initiation codon of human 25 mcg4 is not preceded by an in-frame stop codon and it is therefore possible that the cDNA described in Figure 1 is a truncated form. However, sequence alignment of human and mouse mcg4 ESTs showed a lower degree of nucleotide conservation prior to the assigned initiation codon, thus supporting the notion that the region represents the 5' UTR (Figure 9). To determine the expression pattern of mcg4, 15 $\mu$ g of the total cellular RNA (RNeasy Mini Kit, 30 Qiagen) from various human cell lines grown in culture were electrophoresed through 1.2% w/v MOPS/formaldehyde gels and blotted onto nylon membranes (Amersham) by capillary transfer using 20 x SSC (Sambrook et al, 1989). Filters were subsequently UV-fixed and hybridised overnight at 65°C to a radiolabelled (<sup>32</sup>P-dCTP) cDNA probe (Church and Gilbert, 1984) for mcg4. After washes in 0.1 x SSC/0.1% w/v SDS at 65°C for 1 hour, the filters were air-dried and exposed to X-ray film. This Northern analysis showed that mcg4 is expressed as a 1.6kb message in numerous tissues including breast, ovary, bladder, lung and keratinocytes (Figure 7).

#### **EXAMPLE 7**

A human gene (designated mcg7) was identified and isolated from chromosome 11q13 which 10 encodes a protein that bears striking homology with guanine nucleotide exchange factors (GEFs) from a wide variety of organisms (Fig. 12).

#### **EXAMPLE 8**

15 The composite mcg7 cDNA sequence is at least 2.4kb in length and Figure 13(a) shows a predicted translation product of at least 609 amino acids beginning at methionine 120. An alternative start site due to alternate exon splicing (indicated in lower case) may yield a protein of 671 amino acids starting at methionine 58 (Fig. 13a).

## 20 EXAMPLE 9

An mcg7 homologue from C. elegans has been identified, the product of which is highly conserved with that of MCG7 (Fig. 14). There are several salient features of the protein which have been underlined in Fig. 14 - namely: a guanine nucleotide binding region, a diacylglycerol binding region, and "EF-hand"-calcium binding regions. In addition, there are several potential cAMP, protein kinase C, and casein kinase II phosphorylation sites, as well as a number of potential sites for glycosylation (not indicated).

## **EXAMPLE 10**

contains a cDNA (Acc. no. Y12336) encoding a full-length open reading frame (ORF) for human mcg7 as well as a partial murine mcg7 ORF (Y12339). In addition, the complete genomic sequence of the human mcg7 gene is contained within GenBank entry AC000134.

5 EXAMPLE 11

The best characterised GEFs are members of the family of *ras* oncoproteins, which play a pivotal role in signal transduction and when mutated are responsible for tumour development. A variety of therapeutic regimes for cancer treatment have been designed to specifically interfere with the 10 *ras* signalling pathways. There is potential, therefore that the product of *mcg7* could also be a target for such clinical strategies.

#### **EXAMPLE 12**

- The nucleotide sequence for mcg7 cDNA was extended 5' with genomic DNA sequence from Genbank accession number AC000134 (positions 1-321) and analysed for additional coding sequence 5' to the putative initiation codon (nt 681-683) (Fig. 16). An additional in-frame ATG occurs at position nt 495-497 when the alternatively splice exon (position nt 504-609) is present (also shown in Fig. 13(a)). This closely matches the Kozak consensus. When this exon is absent, then the ATG is not in-frame and other possible initiation codons are absent (resulting translation shown in lower case lettering) (also shown in Fig. 13(b)). Further evidence that the initiation codon at position nt 681-683 is the true initiation site is given in Figure 15.
- Alignment of human and a partial murine mcg7 cDNA sequences is shown in Figure 15. The putative initiation codon is at position nt 360-362. Both murine ESTs appear to have an upstream in-frame stop codon at position nt 326-328, downstream of the differentially spliced exon and the sequence alignment thus suggests that this region represents the 5' UTR of mcg7.

Furthermore, similarity with the *C. elegans* homologue strongly suggest that the ATG codon at position nt 360-362 encodes the N-terminus of MCG7.

## **EXAMPLE 13**

Figure 17 shows data from experiments indicating that a truncated version of MCG7 when expressed as a GST fusion protein (construct B in Fig. 18) can function as a Ras-guanine nucleotide exchange factor. In brief, Ras (unprocessed and as a GST fusion protein) is loaded with <sup>3</sup>H-GDP then incubated in the presence of excess cold GTP ± GST-MCG7. Full details of this assay can be found in Porfiri et al.

## **EXAMPLE 14**

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Nucleotide sequence data generated from cosmid clone cSRL-20h12 with the T7 primer (Promega, and Applied Biosystems Incorporated dye terminator sequencing kit) were aligned to the GenBank Expressed Sequence Tag (EST) database using the program BLASTN (Altschul et al, 1990) and was found to match GenBank entries T78563 (clone 113434) TO9103 (clone HIBBP12) and AA035643 (clone 471819). EST clones 113434 and 471819 were obtained from Genome Systems Inc. and these DNAs were sequenced on both strands with gene-specific primers (Table 5) to generate the cDNA sequence of mcg7 shown in Figures 13(a) and (b).

The cDNA sequence of *mcg*7 was translated in all possible reading frames and compared to the GenBank non-redundant protein database using the program BLASTX (Altschul *et al*, 1990) and the coding region was assigned on the basis of showing homology to the *C. elegans* protein F25B3.3 (Figure 14). The *mcg*7 cDNA composite was suspected to contain a single nucleotide error that originated from clone 471819 and the correct nucleotide sequence was, therefore, sought by reverse transcription-polymerase chain reaction (RT-PCR) of the cDNA fragment from a human cDNA pool. Total RNA was extracted from a human lymphoblastoid cell line using an RNeasy Mini Kit (Qiagen). cDNA synthesis was conducted with the reverse transcriptase Superscript II RNaseH- (GIBCO, BRL) and random hexamers using the procedure recommended by the manufacturer (GIBCO, BRL). One fortieth of the cDNA mix was subjected to 35 cycles of PCR using the following cycling conditions: 94°C for 30 seconds, 58°C for 30 seconds and 72°C for 90 seconds. The 50μl reaction mix consisted of 1x reaction buffer

(Dade Scientific), 2mM dNTP mix, 20pmol of primers (see Table 6) MCG7UF (within the

variably spliced exon of Figure 13(b), between nucleotide positions 184-201) and SGCADRV2 (between nucleotide positions 866-846 of Figure 13(a)) and 10 units of Dynazyme (Dade Scientific). The resulting PCR product was cloned into the pGEM-T vector (Promega) using standard methodology and sequenced using gene-specific primers. The correct nucleotide sequence of mcg7 (as shown in Figure 13(a)) matches that of the recently release GenBank entry Y12336. A partial mouse mcg7 cDNA sequence can also be found in GenBank entry Y12339.

#### **EXAMPLE 15**

The coding sequence of *mcg*7 was cloned into vectors for expression in both bacterial and mammalian cells. In addition to the full-length constructs, the deletion constructs shown in Figure 18 were designed to retain the guanine nucleotide exchange (GEF) domain. For prokaryotic expression, the *mcg*7 coding region was inserted downstream of and in-frame with the Sj26 cassette of the pGEX (Pharmacia) series of vectors (Smith and Johnson, 1988) using standard cloning techniques (Sambrook *et al*, 1989). For mammalian expression, the *mcg*7 coding sequence was first *myc*-tagged at the N-terminus and then ligated into the expression vector pc Exv-n using standard cloning techniques. Ligation junctions of the constructs were sequences as the cloning strategies inadvertently changed or introduced additional amino acids as shown below.

20

Construct (A): EST clone 113434 was digested with ApaI (Figure 13(a), nucleotide positions 1022 to >2416 (within the vector)), blunt-ended with T4 DNA polymerase according to the specifications of the manufacturer (New England Biolab) and ligated into the SmaI site of pGEX-3X.

25

Sequence of the pGEX and mcg7 (underlined) junction:

pGEX-3X

mcg7 (1022)

Sj26 ... GGG ATC CCC CTG GTC [SEQ ID NO:19]

additional amino acids Gly Ile Pro

30

Construct (B): EST clone 113434 was digested with EcoRI (Figure 13(a), nucleotide

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positions <695 (within the vector) to 1711) and ligated into the EcoRI site of pGEX-1.

Sequence of the pGEX and mcg7 (underlined) junction:

pGEX-1

mcg7 (695)

5 Sj26 ... GAA TTC GGC ACG AG<u>C CGA CGG</u> [SEQ ID NO:20] additional amino acids Glu Phe Gly Thr Ser

Construct (C): full-length mcg7: The pGEM-T clone containing the 5' end of the mcg7 coding region was digested with ApaI (subsequently blunt-ended with T4 DNA polymerase) and BstXI to liberate the fragment between nucleotide positions 336 and 830 of Figure 13(a). Clone 113434 was digested with BstXI and HindIII (vector derived) to liberate a fragment between nucleotide positions 830 > and 2416 (vector derived) of Figure 13(a). A pGEM-11zf vector (Promega) containing the myc-tag was digested with ApaI (subsequently blunt-ended with T4 DNA polymerase) and HindIII, and ligated with the 2 inserts described above.

15

Sequence of the myc-tag/mcg7 junction [SEQ ID NOs:21/22]:

The myc-tagged full-length mcg7 insert in pGEM-11zf was then excised with SacI and HindIII (both vector derived) and directionally cloned into the mammalian expression vector pEXV 25 (Beranger et al, 1994).

Construct (D): Construct (C) in pGEM-11zf was sequentially digested with *Hind*III (this site was subsequently blunt-ended with T4 DNA polymerase) then *BamHI*, and ligated into pGEX-2T digested with *BamHI* and *SmaI*. Digestion with *BamHI*, and ligated into pGEX-2T digested with *BamHI* and *SmaI*. Digestion with *BamHI* removed the *myc*-tag of Construct (C).

Sequence of the pGEX and mcg7 [SEQ ID NO:23/24] (underlined) junction:

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pGEX-2 BamHI mcg7 (337)

Sj26 ... gga tcc GCA GCC CAC CCC GCG CCG GCC ATG

Gly Ser Ala Ala His Pro Ala Pro Ala Ala Met

-----additional amino acids------

5

#### **EXAMPLE 16**

Overnight bacterial cultures containing the pGEX plasmid were used to inoculate 500ml of Luria Broth media containing  $50\mu$ g/ml ampicillin. The cultures were grown to an OD of ~0.8 and then 10 induced with 1mM of IPTG for up to 3 hours at 37°C. The bacteria were pelleted and resuspended in 15 ml of STE buffer (10mM Tris pH 8.0, 150 mM NaCl and 1mM EDTA) with 1 mg/ml lysozyme. The mixture was left on ice for more than 1 hour and subsequent steps were performed at 4°C. Protease inhibitors aprotinin, pepstatin and leupeptin were added at final concentrations of 25µg/ml, prior to the addition of Triton-X-100 (2% v/v final) and n-laurovl 15 sarcosine (1.5% w/v final). The lysate was sonicated for ~1 minute and pelleted at 14,000 x g for 15 minutes. 100  $\mu$ l of 50% w/v glutathione-sephadex bead slurry (in PBS) was added per ml of supernatant. Following a 30 minute incubation at 4°C, the beads were washed three times with NETN (20mM Tris-HC1 pH 8.0, 100mM NaCl, 1mM EDTA, 0.5% NP40), once with NETN-HS (equivalent to NETN but with 1M NaCl), and once in NETN. The bound protein 20 was directly analysed by SDS-polyacrylamide gel electrophoresis (PAGE) as described below or the bound protein was eluted from the beads with the following elution buffer (50mM Tris pH 8.0, 150mM NaCl, 5mM MgCl<sub>2</sub>, 1mM DTT, 10mM reduced glutathione) for use in GDP release assays.

25

#### **EXAMPLE 17**

Twenty microlitres of GST-sepharose-bound MCG7 were added to an equal volume of 2 x 30 sample loading dye (100mM Tris pH6.8, 2% v/v mercaptoethanol, 4% w/v SDS, 0.2% w/v bromophenol blue, 20% v/v glycerol), boiled for 5 min and loaded onto a 7.5% w/v SDS-PAGE gel (Sambrook et al, 1989). The Coomassie brilliant blue stained gel (Sambrook et al, 1989)

typically displayed a protein doublet, running between 87-95 kDa consisting of the MCG7-GST fusion and a slightly smaller, co-purified contaminating *E. coli* protein of ~105kDa. The calculated molecular weight of full-length MCG7 is 77.5 kDa (Construct (D)) and the GST component has a molecular weight of 26kDa, hence, the recombinant protein runs slightly smaller than predicted. A Western blot of the same gel probed with anti-GST antibody yields an MCG7-specific band at the same position as that of the stained gel.

#### **EXAMPLE 18**

10 Assumptions: (a) GST-Ras molecular weight = 50 kD; (b) Concentration of GST-Ras solution = 1mg/ml = 20μM; (c) [³H]-GDP is 1mCi/ml and 13.3Ci/mmol, therefore [ H]-GDP concentration = 75 μM and 1pmol [³H]-GDP=15,466 cpm; (d) Elution buffer = Buffer E = 20 mM Tris-Cl, pH7.5; 50mM NaCl; 5mM MgCl<sub>2</sub>; 1mM DTT (added just before use). Buffer E + BSA= Buffer E+1mg/ml BSA (added just before use).

15

Mix together, in the following order and mix well after each addition:

10μl (=10μg) GST-Ras (@1mg/ml in Buffer E), 463μl Buffer E + BSA, 7μl [³H]-GDP, 10ml 490 μM EDTA. Incubate @ RT for 10 min. Add 10μl 0.5 M MgCl<sub>2</sub> and mix well. Incubate @ RT for 10 min. Place on ice. During the first incubation the excess EDTA concentration is 20 5mM, during the second incubation the excess Mg concentration is 5mM. The [³H]-GDP concentration is 1μM and the final concentration of GST-Ras is 400nM. Thus 20ml of the final mix will contain 8pmol of GST-Ras protein. Specific activity of GDP is 15,446 cpm/pmol x (1/1.4) = 11,047 cpm/pmol.

25

#### **EXAMPLE 19**

Exchange Ras with labelled GDP as above. Add unlabelled GTP (stock = 100mM, pH7) to 1 mM. Adjust Mg concentration by adding 5μl 0.5 EDTA to labelled Ras, 5μl 0.5M EDTA to 500μl MCG7, and 5μl 0.5M EDTA to 500μl Buffer E + BSA. On ice set up microfuge tubes with 40μl Ras-GDP (in triplicate) with 40μl MCG7 or Buffer E + BSA (control). Transfer tubes to heat block @ 25°C and incubate for 10, 20 or 30 min. Stop exchange reactions with 1ml of

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ice cold buffer E and place on ice. Pre-soak nitrocellulose filters, pore size 45μm, in Buffer E. Assemble the vacuum manifold apparatus (Millipore) with wet filters and plug the wells with rubber bunds. Switch on the vacuum pump. Remove the first plug, aliquot the sample and once it has been sucked through, wash the filter with 10ml of ice cold Buffer E. Remove next plug etc and continue round the manifold. Take manifold apart. Pin the filters to a pin board reserved for [³H]. Air dry. Take up in 4ml scintillation fluid and count. These studies have been carried out with a truncated MCG7-GST fusion protein (amino acids 341 of Figure 13a to stop encoded within construct B).

10

#### **EXAMPLE 20**

A human gene was identified from chromosome 11q13 that encodes a new member of the DnaJ family of proteins (designated MCG18). This gene (mcg18) is expressed as an ~1.4kb mRNA (Fig. 28) and is predicted to encode a 241 amino acid product (Fig. 19).

15

#### **EXAMPLE 21**

MCG18 has partial homology to E. coli dnaJ and other human DnaJ family members in that it contains the J domain (Fig. 20).

20

#### **EXAMPLE 22**

MCG18 has greatest homology to functionally undefined proteins from *C. elegans* (Fig. 21) and *S. pombe* (Fig. 22) that also feature the J domain but maintain sequence similarity through the central and C-terminal regions of the proteins.

#### **EXAMPLE 23**

The J domain is proposed to mediate interaction with heat shock protein (Hsp70) 70 and consist 30 of some 70 amino acids, frequently located at the N-terminus of the protein. One of these proteins, tumorous imaginal discs (Tid58) from *Drosophila virilis* (Fig. 23) functions as a

tumour suppressor.

#### **EXAMPLE 24**

5 A comparison of homology between MCG18 and human DnaJ proteins HDJ-2/H5DJ, HDJ-1/HSP40 and HSJ1 is shown in Fig. 24.

#### **EXAMPLE 25**

- During the sequence characterisation of the VRF/VEGFB promoter region on cosmid CLGW4 [Grimmond et al, 1996], which maps to chromosome 11q13 the inventors identified a sequence that exactly matched numerous human and mouse expressed sequence tags (ESTs) in the EST database from a gene which we designated mcg18. EST clones for human (GenBank accession number T69741, clone 108172; accession number H40901, clone 177008) and mouse mcg18
  (accession number W34884, clone 350966; accession number W64183, clone 385535) were obtained from Genome Systems Inc. and sequenced with the gene-specific primers shown in Table 7. The EST clones listed in Table 8 were also utilised in generating the full-length coding sequence for human (Figure 19) and mouse (Figure 25) mcg18. The EST database also contained mcg18 cDNA entries that were alternately (or partially) spliced, and in order to understand their ability to encode new polypeptides, the gene structure of mcg18 was determined by sequencing human and mouse genomic templates with gene-specific primers.
- Genomic fragments containing the human [Grimmond et al, 1996] and murine genes [Townson et al, 1996] have been previously reported. Cosmid CLGW4 contains the entire human gene and λ121 contains the entire mouse gene, as determined by direct sequencing of the templates with the oligonucleotides listed in Table 7. Plasmids containing sub-fragments of λ121 and cosmid CLGW4 were prepared using plasmid purification kits (Qiagen) and sequenced as described previously [Grimmond et al, 1996; Townson et al, 1996] using primers designed against cDNA and genomic sequences. The BLAST suite of programs [Altschul et al, 1990] was used to compare the sequence data against the nucleotide and protein databases at the National Center for Biotechnology Information (http//www.ncbi.nih.gov.nlm). The sequence

data were compiled using MacVector 4.2.1 software (IBI-Kodak). ClustalW sequence alignments [Thompson *et al*, 1994] were conducted using the Australian National Genome Information Service computer faculty at the University of Sydney, Australia.

5 The cDNA sequence of human mcg18 (Figure 19) was translated in all possible reading frames and compared to the GenBank non-redundant protein database using the program BLASTX [Altschul et al, 1990] and the coding region was identified on the basis of showing homology to the DnaJ family of proteins (Figure 20). The DnaJ domain is encoded within the longest open reading frame and the assigned initiation codon is preceded by an in-frame stop codon (Figure 27). Similar database search results were obtained for the mouse mcg18 cDNA, and the alignment of human and mouse protein sequences is shown in Figure 26. MCG18 has greatest homology to gene products from C. elegans (Figure 21) and S. pombe (Figure 22). Although it shares a similar J-domain, MCG18 does not contain other domains described for the tumour suppressor gene from D. virilis (Figure 23), nor is it a homologue of other reported human J-domain-containing proteins (Figure 24).

To determine the expression pattern of mcg18,  $15\mu g$  of total cellular RNA (RNeasy Mini Kit, Qiagen) from various human cell lines grown in culture were electrophoresed through 1.2% MOPS/formaldehyde gels and blotted onto nylon membranes (Amersham) by capillary transfer using 20 x SSC (Sambrook et al, 1986). Filters were subsequently UV-fixed and hybridised overnight at 65°C to a radiolabelled ( $^{32}P$ -dCTP) cDNA probe (Church and Gilbert, 1984) for mcg18. After washes in 0.1 x SSC/0.1% w/v SDS for 65°C for 1 hour, the filters were air-dried and exposed to X-ray film. This Northern analysis showed that mcg18 is expressed as a 1.4kb message in numerous tissues including breast, ovary, bladder, lung and keratinocytes (Figure 28).

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# TABLE 4

# ESTs matching mcg4

accession number	seq. run organism	score	E value	N
gb AA399110 AA399110		1136	4.0e-168	2
gb N39612 N39612	yy51g06.s1 Homo sapiens cDNA clone 2	1521	5.3e-168	4
gb AA514406 AA514406	nf57d01.s1 NCI_CGAP_Co3 Homo sapiens	931	5.5e-166	3
gb   AAS44946   AAS44946	vk38e02.rl Soares mouse mammary glan	1207	8.4e-164	2
gb AA450076 AA450076	zx42a04.sl Soares total fetus Nb2HF8	691	2.3e-160	4
gb AA535731 AA535731		796	3.5e-158	4
gb W79710 W79710	zd86f01.rl Soares fetal heart NbHH19	1644	1.1e-157	4
gb AA503531 AA503531	· · · · · · · · · · · · · · · · · · ·	736	4.0e-156	4
gb AA450132 AA450132		1955	3.9e-155	1
gb AA398068 AA398068		1315	5.4e-148	2
gb   W60405   W60405	zd29h08.rl Soares fetal heart NbHH19	1022	1.8e-139	4
gb W81382 W81382	zd86f01.sl Soares fetal heart NbHH19	605	3.5e-125	S
gb AA047617 AA047617		922	4.6e-125	2
gb AA282175 AA282175		1577	2.0e-123	1
gb AA242159 AA242159	my30d04.rl Barstead mouse pooled org	866	7.7e-117	2
gb AA068680 AA068680	mm61a05.rl Stratagene mouse embryoni	1280	1.6e-98	1
gb w46766 w46766	zc36b07.sl Soares senescent fibrobla	506	9.6e-92	3
gb N93704 N93704	zb51c04.sl Soares fetal lung NbHL19W	584	9.0 <b>e-</b> 91	4
gb AA155210 AA155210	mr98e01.rl Stratagene mouse embryoni	840	7.6e-87	2
95 AA366022 AA366022		1077	2.4e-81	1
gb AA037691 AA037691	zk34h12.sl Soares pregnant uterus Nb	949	2.1e-80	2
gb W35374 W35374	zc07h03.sl Soares parathyroid tumor	1016	3.le-76	1
dbj/c00696/c00696	HUMGS0008251, Human Gene Signature,	1009	1.2e-75	1
gb[T98249]T98249	ye59a07.sl Homo sapiens cDNA clone 1	998	6.7e-75	1
gb   W21588   W21588	zb51c04.rl Soares fetal lung NbHL19W	484	1.le-69	4
ды нз2171 нз2171	EST107015 Rattus sp. cDNA 5' end.	828	1.1e-60	1
gb AA108092 AA108092	mm89e06.rl Stratagene mouse embryoni	782	1.3e-60	2
gb AA017857 AA017857	mh44d10.rl Soares mouse placenta 4Nb	665	2.5e-60	2
gb   AA037690   AA037690	zk34h12.rl Soares pregnant uterus Nb	540	9.4e-53	2
	nj07b11.s1 NCI_CGAP_Pr22 Homo sapien	535	5.4e-48	2
gb N46760 N46760	yy51g06.rl Homo sapiens cDNA clone 2	665	9.5e-47	1
gb w23584 w23584	zc71d03.sl Soares fetal heart NbHH19	457	1.8e-44	2
gb W42214 W42214	mc69h09.rl Soares mouse embryo NbME1	460	1.3e-38	3
gb AA244877 AA244877		429	2.9e-25	1
gb w32939 w32939	zc07h03.rl Soares parathyroid tumor	320	4.8e-18	1

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TABLE 5
ESTs matching AA074703 (mcg4-related cDNA)

Database: Non-redundant Database of GenBank EST Division
1,222,625 sequences; 449,352,662 total letters.

Smallest

Sum

			High	Probabili	ty
Sequences producing H	igh-scoring	Segment Pairs:	Score	P (N)	N
accession number	seq. run	organism	score	E value	N
gb AA074703 AA074703	zm76g07.rl	Stratagene neuroepitheli	2071	4.0e-167	1
gb AA068680 AA068680	mm61a05.rl	Stratagene mouse embryon	1270	4.4e-145	4
gb AA134788 AA134788	zm81g02.r1	Stratagene neuroepitheli	946	1.3e-144	5
gb AA399110 AA399110	zt89e06.sl	Soares testis NHT Homo s	520	8.7e-119	6
gb N39612 N39612	yy51g06.s1	Homo sapiens cDNA clone	582	9.6e-110	7
gb AA282175 AA282175	zt02d03.s1	NCI_CGAP_GCB1 Homo sapie	771	9.4e-80	3
gb W81382 W81382	zd86f01.s1	Soares fetal heart NbHHl	329	1.6e-75	6
gb AA544946 AA544946	vk38e02.rl	Soares mouse mammary gla	644	9.6e-63	2
gb w35374 w35374	zc07h03.s1	Soares parathyroid tumor	294	4.5e-42	4
gb W57106 W57106	md57c12.r1	Soares mouse embryo NbME	394	1.9e-30	2
gb AA244877 AA244877	mx25a04.rl	Soares mouse NML Mus mus	162	2.1e-27	4
gb AA017857 AA017857	mh44d10.r1	Soares mouse placenta 4N	230	3.7e-23	3
gb AA531006 AA531006	nj07b11.s1	NCI_CGAP_Pr22 Homo sapie	139	2.3e-19	3
gb H32171 H32171	EST107015 R	Nattus sp. cDNA 5' end.	207	2.6e-10	2
gb W79710 W79710	zd86f01.r1	Soares fetal heart NbHHl	157	0.0073	1

TABLE 6
mcg7-specific oligonucleotides

5	name	sequence (5' to 3')	SEQ ID NOs.
	M1044R	GGA CAA AGT GTG TGA TGA ACC	SEQ ID NO:25
	MCG7-GEF-REV2	CTC ATC CTC CGT CTG ATA CTG	SEQ ID NO:26
	M7R	GTA GAT GTG GAT CAG CTT GG	SEQ ID NO:27
	MCG7 CA FOR	AGG TGG AGA ATG GTC AAGG	SEQ ID NO:28
10	MCG7-GEF-REV	GTC ATA GTC TGT CTC CTA CT	SEQ ID NO:29
	MCG7 GEF FOR	ACA TAG ACA GCG TGC CTA CC	SEQ ID NO:30
	MCG7-PKC-REV	TAC AAC CTT AGG GAC ACC AG	SEQ ID NO:31
	MCG7-PKC-FOR	TGC TGA GCC TGC TCA CGG TG	SEQ ID NO:32
	T09103F	CAA GTG AAC AGC ACG TCC	SEQ ID NO:33
15	M7F	GAC TAT CTC AAG GAC CAG CTG	SEQ ID NO:34
	MCG7UF	GGT TCG GTC CGA GCC CGG	SEQ ID NO:35
	SGCADRV2	GGA GCG ATA CTC CAA GTA GGT	SEQ ID NO:36

TABLE 7
mcg18-SPECIFIC OLIGONUCLEOTIDES

	name	sequence 5' to 3'							
5	HVESTF	AGC GGG CCA GGC CCC TTC [SEQ ID NO:37]							
	HV195F	CAT CCT GGT CCA ATG CGC TC [SEQ ID NO:38]							
	HV387F2	GCA CTG AGG AAG TTA AAC GAG C [SEQ ID NO:39]							
	HV408R	GCT CGT TTA ACT TCC TCA GTG C [SEQ ID NO:40]							
	EXONIREV	GCT CAG CTC CAC AAA GCG GCT [SEQ ID NO:41]							
10	HVEST426F	ACC AGC TCC GCT CAG GTA G [SEQ ID NO:42]							
	HVEST623R	TCC AGG AGC TGT GTG TTT GG [SEQ ID NO:43]							
	SGVESTF3	CCA GTT TCA CAG CGT GAG G [SEQ ID NO:44]							
	HVEST631R	CAG CAT GAG GAG GCA G [SEQ ID NO:45]							

TABLE 8  ${\tt EST\ CLONE\ SEQUENCES\ USED\ TO\ GENERATE\ HUMAN\ AND\ MOUSE}$   ${\tt mcg18\ cDNA\ SEQUENCE\ COMPOSITES}$ 

EST clone number	organism	GenBank accession number
1g2815	human	D45683
001-T2-18	human	F17225
273748	human	N37043
177008	human	H40901 and H40939
258011	human	N30776
276887 -	human	N44004
108172	human	T69741
307529	human	W21083 and W32579
342027	human	W60283
354288	mouse	W44038
350966	mouse	W348844
426261	mouse	AA002868
368185	mouse	W53911
385535	mouse	W64183
404472	mouse	W82959
406437	mouse	W83482

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  Molecular and Cellular Biology 14: 744-758.
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## SEQUENCE LISTING

## (1) GENERAL INFORMATION:

(i) APPLICANT: (OTHER THAN US):

The Council of The Queensland Institute of

Medical Research

(US ONLY): HAYWARD Nicholas, SILINS Ginters, GRIMMOND Sean, GARTSIDE Michael and HANCOCK, John

- (ii) TITLE OF INVENTION: A NOVEL GENE AND USES THEREFOR
- (iii) NUMBER OF SEQUENCES: 45
- (iv) CORRESPONDENCE ADDRESS:
  - (A) ADDRESSEE: DAVIES COLLISON CAVE
  - (B) STREET: 1 LITTLE COLLINS STREET
  - (C) CITY: MELBOURNE
  - (D) STATE: VICTORIA
  - (E) COUNTRY: AUSTRALIA
  - (F) ZIP: 3000
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Floppy disk
  - (B) COMPUTER: IBM PC compatible
  - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
  - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: PCT INTERNATIONAL
  - (B) FILING DATE: 22-MAY-1998
  - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER: PO6973
  - (B) FILING DATE: 23-MAY-1997
  - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER: PO6974
  - (B) FILING DATE: 23-MAY-1997
  - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER: PO6972
  - (B) FILING DATE: 23-MAY-1997

## (C) CLASSIFICATION:

## (vii) PRIOR APPLICATION DATA:

- (A) APPLICATION NUMBER: PP1459
- (B) FILING DATE: 22-JAN-1998
- (C) CLASSIFICATION:

## (vii) PRIOR APPLICATION DATA:

- (A) APPLICATION NUMBER: PP1460
- (B) FILING DATE: 22-JAN-1998
- (C) CLASSIFICATION:

## (vii) PRIOR APPLICATION DATA:

- (A) APPLICATION NUMBER: PP1458
- (B) FILING DATE: 22-JAN-1998
- (C) CLASSIFICATION:

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- (C) TELEX: AA 31787

389

(2) INFO	DRMATION	FOR	SEQ	ID	NO:	: 1	:
----------	----------	-----	-----	----	-----	-----	---

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 8 amino acids
    (B) TYPE: amino acid
    (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Cys Xaa Xaa Cys Xaa Gly Xaa Gly

#### (2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
  (A) LENGTH: 1242 base pairs

  - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (ix) FEATURE:

105

- (A) NAME/KEY: CDS
- (B) LOCATION: 30..959
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

	(111	, 55,	20141		DOCK.		JIV	JEQ .	ID IN	J: Z:						
TCA	GTAA	ACA (	CAGA	GACT(	GG G(	GATC	GATC		GGG Gly							53
AGA Arg	AAG Lys 10	GTG Val	ACC Thr	AAC Asn	CTG Leu	TTC Phe 15	TGC Cys	TTC Phe	GAA Glu	CAT His	CGG Arg 20	GTC Val	AAC Asn	GTC Val	TGC Cys	101
GAG Glu 25	CAC His	TGC Cys	CTG Leu	GTA Val	GCC Ala 30	AAT Asn	CAC His	GCC Ala	AAG Lys	TGC Cys 35	ATC Ile	GTC Val	CAG Gln	TCC Ser	TAC Tyr 40	149
CTG Leu	CAA Gln	TGG Trp	CTC Leu	CAA Gln 45	GAT Asp	AGC Ser	GAC Asp	TAC Tyr	AAC Asn 50	CCC Pro	AAT Asn	TGC Cys	CGC Arg	CTG Leu 55	TGC Cys	197
AAC Asn	ATA Ile	CCC Pro	CTG Leu 60	GCC Ala	AGC Ser	CGA Arg	GAG Glu	ACG Thr 65	ACC Thr	CGC Arg	CTT Leu	GTC Val	TGC Cys 70	TAT Tyr	GAT Asp	245
CTC Leu	TTT Phe	CAC His 75	TGG Trp	GCC Ala	TGC Cys	CTC Leu	AAT Asn 80	GAA Glu	CGT Arg	GCT Ala	GCC Ala	CAG Gln 85	CTA Leu	CCC Pro	CGA Arg	293
AAC Asn	ACG Thr 90	GCA Ala	CCT Pro	GCC Ala	GGC Gly	TAT Tyr 95	CAG Gln	TGC Cys	CCC Pro	AGC Ser	TGC Cys 100	AAT Asn	GGC Gly	CCC Pro	ATC Ile	341

TTC CCC CCA ACC AAC CTG GCT GGC CCC GTG GCC TCC GCA CTG AGA GAG

Phe Pro Pro Thr Asn Leu Ala Gly Pro Val Ala Ser Ala Leu Arg Glu

115

														Pro 135		437
ATC Ile	GAT Asp	GAG Glu	GTG Val 140	GTG Val	AGC Ser	CCA Pro	GAG Glu	CCC Pro 145	GAG Glu	CCC Pro	CTC Leu	AAC Asn	ACG Thr 150	TCT Ser	GAC Asp	485
														CCA Pro		533
														GCC Ala		581
CGG Arg 185	CCC Pro	CCA Pro	GCT Ala	TCC Ser	CCA Pro 190	GGC Gly	CGG Arg	CCC Pro	GAG Glu	CAG Gln 195	CAC His	ACA Thr	GTG Val	ATC Ile	CAC His 200	629
ATG Met	GGC Gly	AAT Asn	CCT Pro	GAG Glu 205	CCC Pro	TTG Leu	ACT Thr	CAC His	GCC Ala 210	CCT Pro	AGG Arg	AAG Lys	GTG Val	TAT Tyr 215	GAT Asp	677
ACG Thr	CGG Arg	GAT Asp	GAT Asp 220	GAC Asp	CGG Arg	ACA Thr	CCA Pro	GGC Gly 225	CTC Leu	CAT His	GGA Gly	GAC Asp	TGT Cys 230	GAC Asp	GAT Asp	725
														CTG Leu		773
AGG Arg	AGC Ser 250	CGG Arg	GCT Ala	GGG Gly	TCT Ser	CGG Arg 255	AAG Lys	CGG Arg	CCG Pro	CTG Leu	ACC Thr 260	CTG Leu	CTC Leu	CAG Gln	CGG Arg	821
GCG Ala 265	GGG Gly	CTG Leu	CTG Leu	CTA Leu	CTC Leu 270	TTG Leu	GGA Gly	CTG Leu	CTG Leu	GGC Gly 275	TTC Phe	CTG Leu	GCC Ala	CTC Leu	CTT Leu 280	869
GCC Ala	CTC Leu	ATG Met	TCT Ser	CGC Arg 285	CTA Leu	GGC Gly	CGG Arg	GCC Ala	GCA Ala 290	GCT Ala	GAC Asp	AGC Ser	GAT Asp	CCC Pro 295	AAC Asn	917
CTG Leu	GAC Asp	CCA Pro	CTC Leu 300	ATG Met	AAC Asn	CCT Pro	CAC His	ATC Ile 305	CGC Arg	GTG Val	GGC Gly	CCC Pro	TCC Ser 310	TGA *		962
GCC	CCT	rgc '	rtgt	GCT	AG G	CCAG	CTAC	G GA	rgtgo	GGTT	CTG!	rggao	GGA (	GAGG	CGGGGT	1022
AAT	GGGG	AGG (	CTGA	GGC	AC C'	rctro	CACTO	G CC	CCTC	rccc	TCA	AGCC	raa (	GACAC	CTAAGA	1082
CCC	CAGA	ccc 2	AAAG	CCAA	GT C	CACC	AGAG'	r GG	CTCG	CAGG	CCA	GCC	rgg A	AGTC	CCCGTG	1142
GGT(	CAAG	CAT '	TTGT	CTTG	AC T	TGCT'	TTCT	c cc	GGT	CTCC	AGC	CTCC	GAC (	CCTC	CGCCCC	1202
ATG	AAGG	AGC '	TGGC	AGGT	GG A	AATA	AACA	A CA	ACTT	гатт						1242

## (2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 310 amino acids
    (B) TYPE: amino acid
    (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Met Gly Leu Cys Lys Cys Pro Lys Arg Lys Val Thr Asn Leu Phe Cys 1 5 10 15 Phe Glu His Arg Val Asn Val Cys Glu His Cys Leu Val Ala Asn His Ala Lys Cys Ile Val Gln Ser Tyr Leu Gln Trp Leu Gln Asp Ser Asp 35 40 45 Tyr Asn Pro Asn Cys Arg Leu Cys Asn Ile Pro Leu Ala Ser Arg Glu Thr Thr Arg Leu Val Cys Tyr Asp Leu Phe His Trp Ala Cys Leu Asn 65 70 75 80 Glu Arg Ala Ala Gln Leu Pro Arg Asn Thr Ala Pro Ala Gly Tyr Gln
85 90 95 Cys Pro Ser Cys Asn Gly Pro Ile Phe Pro Pro Thr Asn Leu Ala Gly Pro Val Ala Ser Ala Leu Arg Glu Lys Leu Ala Thr Val Asn Trp Ala Arg Ala Gly Leu Gly Leu Pro Leu Ile Asp Glu Val Val Ser Pro Glu Pro Glu Pro Leu Asn Thr Ser Asp Phe Ser Asp Trp Ser Ser Phe Asn Ala Ser Ser Thr Pro Gly Pro Glu Glu Val Asp Ser Ala Ser Ala Ala 165 170 175Pro Ala Phe Tyr Ser Arg Ala Pro Arg Pro Pro Ala Ser Pro Gly Arg 180 185 190 Pro Glu Gln His Thr Val Ile His Met Gly Asn Pro Glu Pro Leu Thr His Ala Pro Arg Lys Val Tyr Asp Thr Arg Asp Asp Asp Arg Thr Pro 210 220 Gly Leu His Gly Asp Cys Asp Asp Asp Lys Tyr Arg Arg Arg Pro Ala 225 230 240 Leu Gly Trp Leu Ala Arg Leu Leu Arg Ser Arg Ala Gly Ser Arg Lys Arg Pro Leu Thr Leu Leu Gln Arg Ala Gly Leu Leu Leu Leu Gly Leu Leu Gly Phe Leu Ala Leu Leu Ala Leu Met Ser Arg Leu Gly Arg 280 Ala Ala Ala Asp Ser Asp Pro Asn Leu Asp Pro Leu Met Asn Pro His Ile Arg Val Gly Pro Ser

## (2) INFORMATION FOR SEQ ID NO:4:

305

(i) SEQUENCE CHARACTERISTICS:

PCT/AU98/00380

- 62 -

(A)	LENGTH:	2415	base	pairs
(D)	MUDE	1	:	<u> </u>

- (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: DNA

## (ix) FEATURE:

WO 98/53061

(A) NAME/KEY: CDS
(B) LOCATION: 3..2188

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
CG ATT TCA TTC CTC GCT CCC CAC AGG TCC CTC TCC CCA AAA TAT TCC Ile Ser Phe Leu Ala Pro His Arg Ser Leu Ser Pro Lys Tyr Ser 1 5 10 15	47
CAT CTT GTC CTA GCC CAT CCC CCA GAC TAT CTC AAG GAC CAG CTG TCC His Leu Val Leu Ala His Pro Pro Asp Tyr Leu Lys Asp Gln Leu Ser 20 25 30	95
CCA CGC CCC CGA CCT CCA CTA GGC CTG TGC CAC CCG CTG CCT GCA GGA Pro Arg Pro Arg Pro Pro Leu Gly Leu Cys His Pro Leu Pro Ala Gly 35 40 45	143
AGA CGC CCG GTC CCG GGC CGG GTT AGC CCC ATG GGA ACG CAG CGC CTG Arg Arg Pro Val Pro Gly Arg Val Ser Pro Met Gly Thr Gln Arg Leu 50 55 60	191
TGT GGC CGC GGG ACT CAA GGC TGG CCT GGC TCA AGT GAA CAG CAC GTC Cys Gly Arg Gly Thr Gln Gly Trp Pro Gly Ser Ser Glu Gln His Val 65 70 75	239
CAG GAG GCG ACC TCG TCC GCG GGT TTG CAT TCT GGG GTG GAC GAG CTG Gln Glu Ala Thr Ser Ser Ala Gly Leu His Ser Gly Val Asp Glu Leu 80 90 95	287
GGG GTT CGG TCC GAG CCC GGT GGG AGG CTC CCG GAG CGC AGC CTG GGC Gly Val Arg Ser Glu Pro Gly Gly Arg Leu Pro Glu Arg Ser Leu Gly 100 105 110	335
CCA GCC CAC CCC GCG CCG GCG GCC ATG GCA GGC ACC CTG GAC CTG GAC Pro Ala His Pro Ala Pro Ala Ala Met Ala Gly Thr Leu Asp Leu Asp 115 120 125	383
AAG GGC TGC ACG GTG GAG GAG CTG CTC CGC GGG TGC ATC GAA GCC TTC Lys Gly Cys Thr Val Glu Glu Leu Leu Arg Gly Cys Ile Glu Ala Phe 130 135 140	431
GAT GAC TCC GGG AAG GTG CGG GAC CCG CAG CTG GTG CGC ATG TTC CTC Asp Asp Ser Gly Lys Val Arg Asp Pro Gln Leu Val Arg Met Phe Leu 145 150 155	479
ATG ATG CAC CCC TGG TAC ATC CCC TCC TCT CAG CTG GCG GCC AAG CTG Met Met His Pro Trp Tyr Ile Pro Ser Ser Gln Leu Ala Ala Lys Leu 160 165 170 175	527
CTC CAC ATC TAC CAA CAA TCC CGG AAG GAC AAC TCC AAT TCC CTG CAG Leu His Ile Tyr Gln Gln Ser Arg Lys Asp Asn Ser Asn Ser Leu Gln 180 185	575
GTG AAA ACG TGC CAC CTG GTC AGG TAC TGG ATC TCC GCC TTC CCA GCG Val Lys Thr Cys His Leu Val Arg Tyr Trp Ile Ser Ala Phe Pro Ala 195 200 205	623

WO 98/53061 PCT/AU98/00380

GAG Glu	TTT Phe	GAC Asp 210	TTG Leu	AAC Asn	CCG Pro	GAG Glu	TTG Leu 215	GCT Ala	GAG Glu	CAG Gln	ATC Ile	AAG Lys 220	GAG Glu	CTG Leu	AAG Lys	671
GCT Ala	CTG Leu 225	CTA Leu	GAC Asp	CAA Gln	GAA Glu	GGG Gly 230	AAC Asn	CGA Arg	CGG Arg	CAC His	AGC Ser 235	AGC Ser	CTA Leu	ATC Ile	GAC Asp	719
ATA Ile 240	GAC Asp	AGC Ser	GTC Val	CCT Pro	ACC Thr 245	TAC Tyr	AAG Lys	TGG Trp	AAG Lys	CGG Arg 250	CAG Gln	GTG Val	ACT Thr	CAG Gln	CGG Arg 255	767
AAC Asn	CCT Pro	GTG Val	GGA Gly	CAG Gln 260	AAA Lys	AAG Lys	CGC Arg	AAG Lys	ATG Met 265	TCC Ser	CTG Leu	TTG Leu	TTT Phe	GAC Asp 270	CAC His	815
CTG Leu	GAG Glu	CCC Pro	ATG Met 275	GAG Glu	CTG Leu	GCG Ala	GAG Glu	CAT His 280	CTC Leu	ACC Thr	TAC Tyr	TTG Leu	GAG Glu 285	TAT Tyr	CGC Arg	863
TCC Ser	TTC Phe	TGC Cys 290	AAG Lys	ATC Ile	CTG Leu	TTT Phe	CAG Gln 295	GAC Asp	TAT Tyr	CAC His	AGT Ser	TTC Phe 300	GTG Val	ACT Thr	CAT His	911
GGC Gly	TGC Cys 305	ACT Thr	GTG Val	GAC Asp	AAC Asn	CCC Pro 310	GTC Val	CTG Leu	GAG Glu	CGG Arg	TTC Phe 315	ATC Ile	TCC Ser	CTC Leu	TTC Phe	959
AAC Asn 320	AGC Ser	GTC Val	TCA Ser	CAG Gln	TGG Trp 325	GTG Val	CAG Gln	CTC Leu	ATG Met	ATC Ile 330	CTC Leu	AGC Ser	AAA Lys	CCC Pro	ACA Thr 335	1007
GCC Ala	CCG Pro	CAG Gln	CGG Arg	GCC Ala 340	CTG Leu	GTC Val	ATC Ile	ACA Thr	CAC His 345	TTT Phe	GTC Val	CAC His	GTG Val	GCG Ala 350	GAG Glu	1055
AAG Lys	CTG Leu	CTA Leu	CAG Gln 355	CTG Leu	CAG Gln	AAC Asn	TTC Phe	AAC Asn 360	ACG Thr	CTG Leu	ATG Met	GCA Ala	GTG Val 365	GTC Val	GGG Gly	1103
GGC Gly	CTG Leu	AGC Ser 370	CAC His	AGC Ser	TCC Ser	ATC Ile	TCC Ser 375	CGC Arg	CTC Leu	AAG Lys	GAG Glu	ACC Thr 380	CAC His	AGC Ser	CAC His	1151
GTT Val	AGC Ser 385	CCT Pro	GAG Glu	ACC Thr	ATC Ile	AAG Lys 390	CTC Leu	TGG Trp	GAG Glu	GGT Gly	CTC Leu 395	ACG Thr	GAA Glu	CTA Leu	GTG Val	1199
ACG Thr 400	GCG Ala	ACA Thr	GGC Gly	AAC Asn	TAT Tyr 405	GGC Gly	AAC Asn	TAC Tyr	CGG Arg	CGT Arg 410	CGG Arg	CTG Leu	GCA Ala	GCC Ala	TGT Cys 415	1247
GTG Val	GGC Gly	TTC Phe	CGC Arg	TTC Phe 420	CCG Pro	ATC Ile	CTG Leu	GGT Gly	GTG Val 425	CAC His	CTC Leu	AAG Lys	GAC Asp	CTG Leu 430	GTG Val	1295
GCC Ala	CTG Leu	CAG Gln	CTG Leu 435	GCA Ala	CTG Leu	CCT Pro	GAC Asp	TGG Trp 440	CTG Leu	GAC Asp	CCA Pro	GCC Ala	CGG Arg 445	ACC Thr	CGG Arg	1343
CTC Leu	AAC Asn	GGG Gly 450	GCC Ala	AAG Lys	ATG Met	AAG Lys	CAG Gln 455	CTC Leu	TTT Phe	AGC Ser	ATC Ile	CTG Leu 460	GAG Glu	GAG Glu	CTG Leu	1391
GCC Ala	ATG Met 465	GTG Val	ACC Thr	AGC Ser	CTG Leu	CGG Arg 470	CCA Pro	CCA Pro	GTA Val	CAG Gln	GCC Ala 475	AAC Asn	CCC Pro	GAC Asp	CTG Leu	1439

									CAG Gln							1487
CTG Leu	TAC Tyr	CAG Gln	CTG Leu	TCC Ser 500	CTG Leu	CAG Gln	CGG Arg	GAG Glu	CCG Pro 505	CGC Arg	TCC Ser	AAG Lys	TCC Ser	TCG Ser 510	CCA Pro	1535
									CCC Pro							1583
									CTG Leu							1631
									GTG Val							1679
									GAA Glu							1727
									GAC Asp 585							1775
									TCC Ser							1823
									GTA Val							1871
									CAC His							1919
									CGA Arg							1967
									GTT Val 665							2015
				Leu					CCC Pro							2063
			His					Phe					Pro		AGG Arg	2111
		Ser					Ile		GAG Glu			Val				2159
	Asp					Ile		TTG Leu	TA	ATAG	ATGC	TG T	GGTT	GGAT	С	2208
AAG	GACT	CAT	TCCT	GCCT	TG G	AGAA	AATA	C TT	CAAC	CAGA	GCA	.GGGA	GCC	TGGG	GGTGTC	2268
GGG	GCAG	GAG	GCTG	GGGA	TG G	GGGT	GGGA	T AT	GAGG	GTGG	CAT	GCAG	CTG	AGGG	CAGGGC	2328
										_	_	_				

CAGGGCTGGT G	CCCTAAGG TTGTACAGAC TCTTG	TGAAT ATTTGTATTT	TCCAGATGGA	2388
ATAAAAAGGC CO	GTGTAATT AACCTTC			2415
(2) INFORMAT	ON FOR SEQ ID NO:5:			
(i) SI	QUENCE CHARACTERISTICS: (A) LENGTH: 728 amino aci (B) TYPE: amino acid (D) TOPOLOGY: linear	ds		
(ii) MC	LECULE TYPE: protein			
(xi) SI	QUENCE DESCRIPTION: SEQ I	D NO:5:		

Ile Ser Phe Leu Ala Pro His Arg Ser Leu Ser Pro Lys Tyr Ser His Leu Val Leu Ala His Pro Pro Asp Tyr Leu Lys Asp Gln Leu Ser Pro 20 25 30Arg Pro Arg Pro Pro Leu Gly Leu Cys His Pro Leu Pro Ala Gly Arg Arg Pro Val Pro Gly Arg Val Ser Pro Met Gly Thr Gln Arg Leu Cys 50 60 Gly Arg Gly Thr Gln Gly Trp Pro Gly Ser Ser Glu Gln His Val Gln 65 70 75 80 Glu Ala Thr Ser Ser Ala Gly Leu His Ser Gly Val Asp Glu Leu Gly Val Arg Ser Glu Pro Gly Gly Arg Leu Pro Glu Arg Ser Leu Gly Pro Ala His Pro Ala Pro Ala Ala Met Ala Gly Thr Leu Asp Leu Asp Lys 115 120 125 Gly Cys Thr Val Glu Glu Leu Leu Arg Gly Cys Ile Glu Ala Phe Asp Asp Ser Gly Lys Val Arg Asp Pro Gln Leu Val Arg Met Phe Leu Met Met His Pro Trp Tyr Ile Pro Ser Ser Gln Leu Ala Ala Lys Leu Leu 170 His Ile Tyr Gln Gln Ser Arg Lys Asp Asn Ser Asn Ser Leu Gln Val 185 Lys Thr Cys His Leu Val Arg Tyr Trp Ile Ser Ala Phe Pro Ala Glu 195 200 205 Phe Asp Leu Asn Pro Glu Leu Ala Glu Gln Ile Lys Glu Leu Lys Ala Leu Leu Asp Gln Glu Gly Asn Arg Arg His Ser Ser Leu Ile Asp Ile 230 Asp Ser Val Pro Thr Tyr Lys Trp Lys Arg Gln Val Thr Gln Arg Asn 245 250 250 Pro Val Gly Gln Lys Lys Arg Lys Met Ser Leu Leu Phe Asp His Leu

Glu Pro Met Glu Leu Ala Glu His Leu Thr Tyr Leu Glu Tyr Arg Ser Phe Cys Lys Ile Leu Phe Gln Asp Tyr His Ser Phe Val Thr His Gly Cys Thr Val Asp Asn Pro Val Leu Glu Arg Phe Ile Ser Leu Phe Asn 310 315 Ser Val Ser Gln Trp Val Gln Leu Met Ile Leu Ser Lys Pro Thr Ala Pro Gln Arg Ala Leu Val Ile Thr His Phe Val His Val Ala Glu Lys 345 Leu Leu Gln Leu Gln Asn Phe Asn Thr Leu Met Ala Val Val Gly Gly Leu Ser His Ser Ser Ile Ser Arg Leu Lys Glu Thr His Ser His Val Ser Pro Glu Thr Ile Lys Leu Trp Glu Gly Leu Thr Glu Leu Val Thr Ala Thr Gly Asn Tyr Gly Asn Tyr Arg Arg Arg Leu Ala Ala Cys Val 410 Gly Phe Arg Phe Pro Ile Leu Gly Val His Leu Lys Asp Leu Val Ala Leu Gln Leu Ala Leu Pro Asp Trp Leu Asp Pro Ala Arg Thr Arg Leu Asn Gly Ala Lys Met Lys Gln Leu Phe Ser Ile Leu Glu Glu Leu Ala Met Val Thr Ser Leu Arg Pro Pro Val Gln Ala Asn Pro Asp Leu Leu Ser Leu Leu Thr Val Ser Leu Asp Gln Tyr Gln Thr Glu Asp Glu Leu Tyr Gln Leu Ser Leu Gln Arg Glu Pro Arg Ser Lys Ser Ser Pro Thr Ser Pro Thr Ser Cys Thr Pro Pro Pro Arg Pro Pro Val Leu Glu Glu Trp Thr Ser Ala Ala Lys Pro Lys Leu Asp Gln Ala Leu Val Val Glu His Ile Glu Lys Met Val Glu Ser Val Phe Arg Asn Phe Asp Val Asp Gly Asp Gly His Ile Ser Gln Glu Glu Phe Gln Ile Ile Arg Gly Asn Phe Pro Tyr Leu Ser Ala Phe Gly Asp Leu Asp Gln Asn Gln Asp Gly 'Cys Ile Ser Arg Glu Glu Met Val Ser Tyr Phe Leu Arg Ser Ser Ser Val Leu Gly Gly Arg Met Gly Phe Val His Asn Phe Gln Glu Ser Asn Ser Leu Arg Pro Val Ala Cys Arg His Cys Lys Ala Leu Ile Leu Gly

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625	630	635		640
Ile Tyr Lys Gln Gly 645	Leu Lys Cys	Arg Ala Cys 650	Gly Val Asn Cys 655	
Lys Gln Cys Lys Asp 660		Val Glu Cys 665	Arg Arg Arg Ala 670	Gln
Ser Val Ser Leu Glu 675	Gly Ser Ala : 680	Pro Ser Pro	Ser Pro Met His 685	s Ser
His His His Arg Ala 690	Phe Ser Phe 695		Arg Pro Gly Arg 700	Arg
Gly Ser Arg Pro Pro 705	Glu Ile Arg (	Glu Glu Glu 715	Val Gln Thr Val	. Glu 720
Asp Gly Val Phe Asp 725	Ile His Leu			
(B) TYPE: (C) STRANI	_	pairs		
(ii) MOLECULE TY	PE: DNA			
(ix) FEATURE: (A) NAME/F (B) LOCATI	KEY: CDS ION: 2542083	3		
(xi) SEQUENCE DE	ESCRIPTION: SE	EQ ID NO:6:		
CGATTTCATT CCTCGCTCC	CC CACAGGTCCC	TCTCCCCAAA A	ATATTCCCAT CTTG	TCCTAG 60
CCCATCCCC AGACTATCT	C AAGGACCAGC	TGTCCCCACG (	CCCCGACCT CCAC	TAGGCC 120
TGTGCCACCC GCTGCCTGC	CA GGAAGACGCC	CGGTCCCGGG (	CCGGGTTAGC CCCA	TGGGAA 180
CGGGGTTCGG TCCGAGCCC	G GTGGGAGGCT	CCCGGAGCGC A	AGCCTGGGCC CAGC	CCACCC 240
CGCGCCGGCG GCC ATG G Met A 1	GCA GGC ACC CT Ala Gly Thr Le	TG GAC CTG GA Bu Asp Leu As 5	AC AAG GGC TGC sp Lys Gly Cys 10	ACG 289 Thr
GTG GAG GAG CTG CTC Val Glu Glu Leu Leu 15	CGC GGG TGC A Arg Gly Cys 1 20	ATC GAA GCC 1	TTC GAT GAC TCC Phe Asp Asp Ser 25	GGG 337 Gly
AAG GTG CGG GAC CCG Lys Val Arg Asp Pro 30	CAG CTG GTG C Gln Leu Val A 35	CGC ATG TTC ( Arg Met Phe I	CTC ATG ATG CAC Leu Met Met His 40	CCC 385 Pro
TGG TAC ATC CCC TCC Trp Tyr Ile Pro Ser 45	TCT CAG CTG C Ser Gln Leu A 50	GCG GCC AAG ( Ala Ala Lys 1 55	CTG CTC CAC ATC Leu Leu His Ile	TAC 433 Tyr 60
CAA CAA TCC CGG AAG Gln Gln Ser Arg Lys 65	GAC AAC TCC A Asp Asn Ser A	AAT TCC CTG ( Asn Ser Leu ( 70	CAG GTG AAA ACG Gln Val Lys Thr 75	TGC 481 Cys
CAC CTG GTC AGG TAC	TGG ATC TCC G	GCC TTC CCA (	GCG GAG TTT GAC	TTG 529

529

His	Leu	Val	Arg 80	Tyr	Trp	Ile	Ser	Ala 85	Phe	Pro	Ala	Glu	Phe 90	Asp	Leu	
AAC Asn	CCG Pro	GAG Glu 95	TTG Leu	GCT Ala	GAG Glu	CAG Gln	ATC Ile 100	AAG Lys	GAG Glu	CTG Leu	AAG Lys	GCT Ala 105	CTG Leu	CTA Leu	GAC Asp	577
CAA Gln	GAA Glu 110	GGG Gly	AAC Asn	CGA Arg	CGG Arg	CAC His 115	AGC Ser	AGC Ser	CTA Leu	ATC Ile	GAC Asp 120	ATA Ile	GAC Asp	AGC Ser	GTC Val	625
CCT Pro 125	ACC Thr	TAC Tyr	AAG Lys	TGG Trp	AAG Lys 130	CGG Arg	CAG Gln	GTG Val	ACT Thr	CAG Gln 135	CGG Arg	AAC Asn	CCT Pro	GTG Val	GGA Gly 140	673
CAG Gln	AAA Lys	AAG Lys	CGC Arg	AAG Lys 145	ATG Met	TCC Ser	CTG Leu	TTG Leu	TTT Phe 150	GAC Asp	CAC His	CTG Leu	GAG Glu	CCC Pro 155	ATG Met	721
GAG Glu	CTG Leu	GCG Ala	GAG Glu 160	CAT His	CTC Leu	ACC Thr	TAC Tyr	TTG Leu 165	GAG Glu	TAT Tyr	CGC Arg	TCC Ser	TTC Phe 170	TGC Cys	AAG Lys	769
ATC Ile	CTG Leu	TTT Phe 175	CAG Gln	GAC Asp	TAT Tyr	CAC His	AGT Ser 180	TTC Phe	GTG Val	ACT Thr	CAT His	GGC Gly 185	TGC Cys	ACT Thr	GTG Val	817
GAC Asp	AAC Asn 190	CCC Pro	GTC Val	CTG Leu	GAG Glu	CGG Arg 195	TTC Phe	ATC Ile	TCC Ser	CTC Leu	TTC Phe 200	AAC Asn	AGC Ser	GTC Val	TCA Ser	865
CAG Gln 205	TGG Trp	GTG Val	CAG Gln	CTC Leu	ATG Met 210	ATC Ile	CTC Leu	AGC Ser	AAA Lys	CCC Pro 215	ACA Thr	GCC Ala	CCG Pro	CAG Gln	CGG Arg 220	913
GCC Ala	CTG Leu	GTC Val	ATC Ile	ACA Thr 225	CAC His	TTT Phe	GTC Val	CAC His	GTG Val 230	GCG Ala	GAG Glu	AAG Lys	CTG Leu	CTA Leu 235	CAG Gln	961
CTG Leu	CAG Gln	AAC Asn	TTC Phe 240	AAC Asn	ACG Thr	CTG Leu	ATG Met	GCA Ala 245	GTG Val	GTC Val	GGG Gly	GGC Gly	CTG Leu 250	AGC Ser	CAC His	1009
AGC Ser	TCC Ser	ATC Ile 255	TCC Ser	CGC Arg	CTC Leu	AAG Lys	GAG Glu 260	ACC Thr	CAC His	AGC Ser	CAC His	GTT Val 265	AGC Ser	CCT Pro	GAG Glu	1057
ACC Thr	ATC Ile 270	AAG Lys	CTC Leu	TGG Trp	GAG Glu	GGT Gly 275	CTC Leu	ACG Thr	GAA Glu	CTA Leu	GTG Val 280	ACG Thr	GCG Ala	ACA Thr	GGC Gly	1105
AAC Asn 285	TAT Tyr	GGC Gly	AAC Asn	TAC Tyr	CGG Arg 290	CGT Arg	CGG Arg	CTG Leu	GCA Ala	GCC Ala 295	TGT Cys	GTG Val	GGC Gly	TTC Phe	CGC Arg 300	1153
TTC Phe	CCG Pro	ATC Ile	CTG Leu	GGT Gly 305	GTG Val	CAC His	CTC Leu	AAG Lys	GAC Asp 310	CTG Leu	GTG Val	GCC Ala	CTG Leu	CAG Gln 315	CTG Leu	1201
GCA 'Ala	CTG Leu	CCT Pro	GAC Asp 320	TGG Trp	CTG Leu	GAC Asp	CCA Pro	GCC Ala 325	CGG Arg	ACC Thr	CGG Arg	CTC Leu	AAC Asn 330	GGG Gly	GCC Ala	1249
AAG Lys	ATG Met	AAG Lys 335	CAG Gln	CTC Leu	TTT Phe	AGC Ser	ATC Ile 340	CTG Leu	GAG Glu	GAG Glu	CTG Leu	GCC Ala 345	ATG Met	GTG Val	ACC Thr	1297

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AGC Ser	CTG Leu 350	CGG Arg	CCA Pro	CCA Pro	GTA Val	CAG Gln 355	GCC Ala	AAC Asn	CCC Pro	GAC Asp	CTG Leu 360	CTG Leu	AGC Ser	CTG Leu	CTC Leu	1345
ACG Thr 365	GTG Val	TCT Ser	CTG Leu	GAT Asp	CAG Gln 370	TAT Tyr	CAG Gln	ACG Thr	GAG Glu	GAT Asp 375	GAG Glu	CTG Leu	TAC Tyr	CAG Gln	CTG Leu 380	1393
TCC Ser	CTG Leu	CAG Gln	CGG Arg	GAG Glu 385	CCG Pro	CGC Arg	TCC Ser	AAG Lys	TCC Ser 390	TCG Ser	CCA Pro	ACC Thr	AGC Ser	CCC Pro 395	ACG Thr	1441
AGT Ser	TGC Cys	ACC Thr	CCA Pro 400	CCA Pro	CCC Pro	CGG Arg	CCC Pro	CCG Pro 405	GTA Val	CTG Leu	GAG Glu	GAG Glu	TGG Trp 410	ACC Thr	TCG Ser	1489
GCT Ala	GCC Ala	AAA Lys 415	CCC Pro	AAG Lys	CTG Leu	GAT Asp	CAG Gln 420	GCC Ala	CTC Leu	GTG Val	GTG Val	GAG Glu 425	CAC His	ATC Ile	GAG Glu	1537
AAG Lys	ATG Met 430	GTG Val	GAG Glu	TCT Ser	GTG Val	TTC Phe 435	CGG Arg	AAC Asn	TTT Phe	GAC Asp	GTC Val 440	GAT Asp	GGG Gly	GAT Asp	GGC Gly	1585
CAC His 445	ATC Ile	TCA Ser	CAG Gln	GAA Glu	GAA Glu 450	TTC Phe	CAG Gln	ATC Ile	ATC Ile	CGT Arg 455	GGG Gly	AAC Asn	TTC Phe	CCT Pro	TAC Tyr 460	1633
CTC Leu	AGC Ser	GCC Ala	TTT Phe	GGG Gly 465	GAC Asp	CTC Leu	GAC Asp	CAG Gln	AAC Asn 470	CAG Gln	GAT Asp	GGC Gly	TGC Cys	ATC Ile 475	AGC Ser	1681
AGG Arg	GAG Glu	GAG Glu	ATG Met 480	GTT Val	TCC Ser	TAT Tyr	TTC Phe	CTG Leu 485	CGC Arg	TCC Ser	AGC Ser	TCT Ser	GTG Val 490	TTG Leu	GGG	1729
GGG Gly	CGC Arg	ATG Met 495	GGC Gly	TTC Phe	GTA Val	CAC His	AAC Asn 500	TTC Phe	CAG Gln	GAG Glu	AGC Ser	AAC Asn 505	TCC Ser	TTG Leu	CGC Arg	1777
CCC Pro	GTC Val 510	GCC Ala	TGC Cys	CGC Arg	CAC His	TGC Cys 515	AAA Lys	GCC Ala	CTG Leu	ATC Ile	CTG Leu 520	GGC Gly	ATC Ile	TAC Tyr	AAG Lys	1825
CAG Gln 525	GGC Gly	CTC Leu	AAA Lys	TGC Cys	CGA Arg 530	GCC Ala	TGT Cys	GGA Gly	GTG Val	AAC Asn 535	TGC Cys	CAC His	AAG Lys	CAG Gln	TGC Cys 540	1873
AAG Lys	GAT Asp	CGC Arg	CTG Leu	TCA Ser 545	GTT Val	GAG Glu	TGT Cys	CGG Arg	CGC Arg 550	AGG Arg	GCC Ala	CAG Gln	AGT Ser	GTG Val 555	AGC Ser	1921
CTG Leu	GAG Glu	GGG Gly	TCT Ser 560	GCA Ala	CCC Pro	TCA Ser	CCC Pro	TCA Ser 565	CCC Pro	ATG Met	CAC His	AGC Ser	CAC His 570	CAT His	CAC His	1969
CGC Arg	GCC Ala	TTC Phe 575	AGC Ser	TTC Phe	TCT Ser	CTG Leu	CCC Pro 580	CGC Arg	CCT Pro	GGC Gly	AGG Arg	CGA Arg 585	GGC Gly	TCC Ser	AGG Arg	2017
ĆCT Pro	CCA Pro 590	GAG Glu	ATC Ile	CGT Arg	GAG Glu	GAG Glu 595	GAG Glu	GTA Val	CAG Gln	ACG Thr	GTG Val 600	GAG Glu	GAT Asp	GGG Gly	GTG Val	2065
TTT Phe 605	GAC Asp	ATC Ile	CAC His	TTG Leu	TAA7	AGAT	GC I	rgtgg	GTTGG	SA TO	CAAGO	SACTO	ATI	CCTG	CCT	2120

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TTAACCTTC						2309
GGTTGTACAG	ACTCTTGTGA	ATATTTGTAT	TTTCCAGATG	GAATAAAAAG	GCCCGTGTAA	2300
TGGGGGTGGG	ATATGAGGGT	GGCATGCAGC	TGAGGGCAGG	GCCAGGGCTG	GTGTCCCTAA	2240
TGGAGAAAAT	ACTTCAACCA	GAGCAGGGAG	CCTGGGGGTG	TCGGGGCAGG	AGGCTGGGGA	2180

#### (2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 609 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:7: Met Ala Gly Thr Leu Asp Leu Asp Lys Gly Cys Thr Val Glu Glu Leu Leu Arg Gly Cys Ile Glu Ala Phe Asp Asp Ser Gly Lys Val Arg Asp 20 25 30 Pro Gln Leu Val Arg Met Phe Leu Met Met His Pro Trp Tyr Ile Pro Ser Ser Gln Leu Ala Ala Lys Leu Leu His Ile Tyr Gln Gln Ser Arg Lys Asp Asn Ser Asn Ser Leu Gln Val Lys Thr Cys His Leu Val Arg Tyr Trp Ile Ser Ala Phe Pro Ala Glu Phe Asp Leu Asn Pro Glu Leu Ala Glu Gln Ile Lys Glu Leu Lys Ala Leu Leu Asp Gln Glu Gly Asn 105 Arg Arg His Ser Ser Leu Ile Asp Ile Asp Ser Val Pro Thr Tyr Lys Trp Lys Arg Gln Val Thr Gln Arg Asn Pro Val Gly Gln Lys Lys Arg Lys Met Ser Leu Leu Phe Asp His Leu Glu Pro Met Glu Leu Ala Glu 155 His Leu Thr Tyr Leu Glu Tyr Arg Ser Phe Cys Lys Ile Leu Phe Gln Asp Tyr His Ser Phe Val Thr His Gly Cys Thr Val Asp Asn Pro Val Leu Glu Arg Phe Ile Ser Leu Phe Asn Ser Val Ser Gln Trp Val Gln Leu Met Ile Leu Ser Lys Pro Thr Ala Pro Gln Arg Ala Leu Val Ile Thr His Phe Val His Val Ala Glu Lys Leu Leu Gln Leu Gln Asn Phe

Asn Thr Leu Met Ala Val Val Gly Gly Leu Ser His Ser Ser Ile Ser

250

255

Arg Leu Lys Glu Thr His Ser His Val Ser Pro Glu Thr Ile Lys Leu Trp Glu Gly Leu Thr Glu Leu Val Thr Ala Thr Gly Asn Tyr Gly Asn Tyr Arg Arg Arg Leu Ala Ala Cys Val Gly Phe Arg Phe Pro Ile Leu Gly Val His Leu Lys Asp Leu Val Ala Leu Gln Leu Ala Leu Pro Asp Trp Leu Asp Pro Ala Arg Thr Arg Leu Asn Gly Ala Lys Met Lys Gln 330 Leu Phe Ser Ile Leu Glu Glu Leu Ala Met Val Thr Ser Leu Arg Pro 345 Pro Val Gln Ala Asn Pro Asp Leu Leu Ser Leu Leu Thr Val Ser Leu Asp Gln Tyr Gln Thr Glu Asp Glu Leu Tyr Gln Leu Ser Leu Gln Arg Glu Pro Arg Ser Lys Ser Ser Pro Thr Ser Pro Thr Ser Cys Thr Pro Pro Pro Arg Pro Pro Val Leu Glu Glu Trp Thr Ser Ala Ala Lys Pro Lys Leu Asp Gln Ala Leu Val Val Glu His Ile Glu Lys Met Val Glu Ser Val Phe Arg Asn Phe Asp Val Asp Gly Asp Gly His Ile Ser Gln Glu Glu Phe Gln Ile Ile Arg Gly Asn Phe Pro Tyr Leu Ser Ala Phe Gly Asp Leu Asp Gln Asn Gln Asp Gly Cys Ile Ser Arg Glu Glu Met Val Ser Tyr Phe Leu Arg Ser Ser Ser Val Leu Gly Gly Arg Met Gly Phe Val His Asn Phe Gln Glu Ser Asn Ser Leu Arg Pro Val Ala Cys 505 Arg His Cys Lys Ala Leu Ile Leu Gly Ile Tyr Lys Gln Gly Leu Lys Cys Arg Ala Cys Gly Val Asn Cys His Lys Gln Cys Lys Asp Arg Leu 535 Ser Val Glu Cys Arg Arg Arg Ala Gln Ser Val Ser Leu Glu Gly Ser 555 Ala Pro Ser Pro Ser Pro Met His Ser His His Arg Ala Phe Ser Phe Ser Leu Pro Arg Pro Gly Arg Arg Gly Ser Arg Pro Pro Glu Ile Arg Glu Glu Glu Val Gln Thr Val Glu Asp Gly Val Phe Asp Ile His 600

(2)	INFORMATION	FOR	SEO	TD	NO · R ·

( i )	SECTIONES	CHARACTERISTICS:
1 1	SECUENCE	CHARACTERISTICS:

- EQUENCE CHARACTERISTICS:

  (A) LENGTH: 832 base pairs

  (B) TYPE: nucleic acid

  (C) STRANDEDNESS: single

  (D) TOPOLOGY: linear

#### (ii) MOLECULE TYPE: DNA

- (ix) FEATURE:
  (A) NAME/KEY: CDS
  (B) LOCATION: 11..733

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

GCC	CGCC	GCC 1	ATG ( Met 1	CCG Pro	CCC Pro	TTA ( Leu )	CTG Leu 5	CCC Pro	CTG Leu	CGC Arg	CTG Leu	TGC ( Cys )	CGG Arg	CTG Leu	TGG Trp	49
CCC Pro	CGC Arg 15	AAC Asn	CCT Pro	CCC Pro	TCC Ser	CGG Arg 20	CTC	CTC Leu	GGA Gly	GCG Ala	GCC Ala 25	GCC Ala	GGG Gly	CAG Gln	CGG Arg	97
TCC Ser 30	AGA Arg	CCC Pro	AGT Ser	ACT Thr	TAT Tyr 35	TAT Tyr	GAA Glu	CTG Leu	TTG Leu	GGG Gly 40	Val	CAT His	CCT Pro	GGT Gly	GCC Ala 45	145
AGC Ser	ACT Thr	GAG Glu	GAA Glu	GTT Val 50	AAA Lys	CGA Arg	GCT Ala	TTC Phe	TTC Phe 55	TCC Ser	AAG Lys	TCC Ser	AAA Lys	GAG Glu 60	CTG Leu	193
CAC His	CCA Pro	GAC Asp	CGG Arg 65	GAC Asp	CCT Pro	GGG Gly	AAC Asn	CCA Pro 70	AGC Ser	CTG Leu	CAC His	AGC Ser	CGC Arg 75	TTT Phe	GTG Val	241
GAG Glu	CTG Leu	AGC Ser 80	GAG Glu	GCA Ala	TAC Tyr	CGT Arg	GTG Val 85	CTC Leu	AGC Ser	CGT Arg	GAG Glu	CAG Gln 90	AGC Ser	CGC Arg	CGC Arg	289
AGC Ser	TAT Tyr 95	GAT Asp	GAC Asp	CAG Gln	CTC Leu	CGC Arg 100	TCA Ser	GGT Gly	AGT Ser	CCC Pro	CCA Pro 105	AAG Lys	TCT Ser	CCA Pro	CGA Arg	337
ACC Thr 110	ACA Thr	GTC Val	CAT His	GAC Asp	AAG Lys 115	TCT Ser	GCC Ala	CAC His	CAA Gln	ACA Thr 120	CAC His	AGC Ser	TCC Ser	TGG Trp	ACA Thr 125	385
CCC Pro	CCC Pro	AAC Asn	GCA Ala	CAG Gln 130	TAC Tyr	TGG Trp	TCC Ser	CAG Gln	TTT Phe 135	CAC His	AGC Ser	GTG Val	AGG Arg	CCA Pro 140	CAG Gln	433
GGG Gly	CCC Pro	CAG Gln	TTG Leu 145	AGG Arg	CAG Gln	CAG Gln	CAA Gln	CAC His 150	AAA Lys	CAA Gln	AAC Asn	AAA Lys	CAA Gln 155	GTG Val	CTG Leu	481
GGG Gly	TAC Tyr	TGC Cys 160	CTC Leu	CTC Leu	CTC Leu	ATG Met	CTG Leu 165	GCG Ala	GGC Gly	ATG Met	GGC Gly	CTG Leu 170	CAC His	TAC Tyr	ATT Ile	529
GCC Ala	TTC Phe 175	AGG Arg	AAG Lys	GTG Val	AAG Lys	CAG Gln 180	ATG Met	CAC His	CTT Leu	AAC Asn	TTC Phe 185	ATG Met	GAT Asp	GAA Glu	AAG Lys	577

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													CGG Arg			625
													GGG Gly			673
													GTG Val 235			721
		GGC Gly 240		TGA *	GGGG	CTC	ACCT	GGAT	GG G	GCCT	CGCAG	T GC	GTTC	CCGC		773
TTTC	CTTC	CT 1	CCCI	GGAC	G GC	CCGC	TCCC	: CGA	AACG	CGC	GCAA	AAAT	GT G	ATTC	GCAG	832

#### (2) INFORMATION FOR SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 241 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9: Met Pro Pro Leu Leu Pro Leu Arg Leu Cys Arg Leu Trp Pro Arg Asn 1 5 10 15 Pro Pro Ser Arg Leu Leu Gly Ala Ala Ala Gly Gln Arg Ser Arg Pro
20 25 30 Ser Thr Tyr Tyr Glu Leu Leu Gly Val His Pro Gly Ala Ser Thr Glu 35 40 45 Glu Val Lys Arg Ala Phe Phe Ser Lys Ser Lys Glu Leu His Pro Asp Arg Asp Pro Gly Asn Pro Ser Leu His Ser Arg Phe Val Glu Leu Ser 65 70 75 80 Glu Ala Tyr Arg Val Leu Ser Arg Glu Gln Ser Arg Arg Ser Tyr Asp 85 90 95 Asp Gln Leu Arg Ser Gly Ser Pro Pro Lys Ser Pro Arg Thr Thr Val His Asp Lys Ser Ala His Gln Thr His Ser Ser Trp Thr Pro Pro Asn Ala Gln Tyr Trp Ser Gln Phe His Ser Val Arg Pro Gln Gly Pro Gln

Leu Arg Gln Gln His Lys Gln Asn Lys Gln Val Leu Gly Tyr Cys

135

Leu Leu Met Leu Ala Gly Met Gly Leu His Tyr Ile Ala Phe Arg 165

Lys Val Lys Gln Met His Leu Asn Phe Met Asp Glu Lys Asp Arg Ile

Ile Thr Ala Phe Tyr Asn Glu Ala Arg Ala Arg Ala Arg Ala Asn Arg

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		195					200					205				
	Ile 210	Leu	Gln	Gln	Glu	Arg 215	Gln	Arg	Leu	Gly	Gln 220	Arg	Gln	Pro	Pro	
Pro 225	Ser	Glu	Pro	Thr	Gln 230	Gly	Pro	Glu	Ile	Val 235	Pro	Arg	Gly	Ala	Gly 240	
Pro																
SEQ	ID N	ios:	10-1	8 25	6-36											
(2)	INFO	ORMAT	NOI	FOR	SEQ	ID 1	NO:7:	:								
	(i)	(E	QUENC ) LE B) TY C) ST O) TC	NGTI PE: RANI	i: 30 nucl	00 ba leic ESS:	ase p acio sino	pairs i	5							
	(ii)	MOI	ECUL	E T	PE:	DNA										
	(ix)		ATURE A) NA B) LC	ME/I			300	0								
	(xi)	) SEC	QUENC	E D	ESCR:	IPTI	ON: 8	SEQ :	ID NO	0:7:						
CGAT	TTC	ATT (	CTC	CTC	CC C	ACAG	GTCC	C TC	rccc	CAAA	ATA:	rtcc	CAT (	CTTG	CCTA	G 60
CCCA	TCC	CCC A	AGACT	'ATC'	rc A	AGGA	CCAG	C TG	rccc	CACG	CCC	CCGA	CT (	CCAC	ragge	C 120
TGTG	CCA	ccc c	CTGC	CTG	CA GO	GAAG	ACGC	C CG	GTCC	CGGG	CCG(	GGTT/		CC CA ro H: 1		175
		GGG Gly 5														223
		CCA Pro														271
		AAG Lys														300
(2)		ORMA'	SEQUI (A (B	ENCE ) LE ) TY	CHA NGTH PE:	RACT : 44 amin		TICS no a id								

(ii) MOLECULE TYPE: protein

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Pro His Gly Asn Gly Val Arg Ser Glu Pro Gly Gly Arg Leu Pro Glu

Arg Ser Leu Gly Pro Ala His Pro Ala Pro Ala Ala Met Ala Gly Thr

Leu Asp Leu Asp Lys Gly Cys Thr Val Glu Glu Leu

- (2) INFORMATION FOR SEQ ID NO:9:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 15 base pairs(B) TYPE: nucleic acid

    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

GGGATCCCCC TGGTC

15

- (2) INFORMATION FOR SEQ ID NO:10:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Asp Val Asp Glu Glu Asp Glu Val Glu Asp Ile Glu Phe

- (2) INFORMATION FOR SEQ ID NO:11:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Asp Val Asp Gly Asp Gly His Ile Ser Gln Glu Glu Phe

- (2) INFORMATION FOR SEQ ID NO:12:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13 amino acids

- (B) TYPE: amino acid(C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Asp His Asp Arg Asp Gly Phe Ile Ser Gln Glu Glu Phe

- (2) INFORMATION FOR SEQ ID NO:13:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Asp Gln Asn Gln Asp Gly Cys Ile Ser Arg Glu Glu Met

- (2) INFORMATION FOR SEQ ID NO:14:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 13 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Asp Val Asp Met Asp Gly Gln Ile Ser Lys Asp Glu Leu

- (2) INFORMATION FOR SEQ ID NO:15:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 37 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

His Phe Val His Val Ala Glu Lys Leu Leu Gln Leu Gln Asn Phe Asn

Thr Leu Met Ala Val Val Gly Gly Leu Ser His Ser Ser Ile Ser Arg

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Leu Lys Glu Thr His 35

- (2) INFORMATION FOR SEQ ID NO:16:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 37 amino acids(B) TYPE: amino acid

    - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Lys Phe Val His Val Ala Lys His Leu Arg Lys Ile Asn Asn Phe Asn

Thr Leu Met Ser Val Val Gly Gly Ile Thr His Ser Ser Val Ala Arg

Leu Ala Lys Thr Tyr 35

- (2) INFORMATION FOR SEQ ID NO:17:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 50 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

His Asn Phe Gln Glu Ser Asn Ser Leu Arg Pro Val Ala Cys Arg His

Cys Lys Ala Leu Ile Leu Gly Ile Tyr Lys Gln Gly Leu Lys Cys Arg 20 25 30

Ala Cys Gly Val Asn Cys His Lys Gln Cys Lys Asp Arg Leu Ser Val

Glu Cys 50

- (2) INFORMATION FOR SEQ ID NO:18:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 50 amino acids

    - (B) TYPE: amino acid (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: Peptide
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

His Asn Phe His Glu Thr Thr Phe Leu Thr Pro Thr Thr Cys Asn His

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	1				5					10					15		
	Суѕ	Asn	Lys	Leu 20	Leu	Trp	Gly	Ile	Leu 25	Arg	Gln	Gly	Phe	Lys 30	Cys	Lys	
	Asp	Cys	Gly 35	Leu	Ala	Val	His	Ser 40	Cys	Cys	Lys	Ser	Asn 45	Ala	Val	Ala	
	Glu	Суs 50															
(2)	INFO	RMAT:	ION I	FOR :	SEQ I	ID N	0:19	:									
	(i)	(A) (B) (C)	LEI TY STI	NGTH PE: 1 RAND	ARACT : 15 nucle EDNES GY: 1	base eic a SS: s	e pa: acid sing:	irs							,		
	(ii)	MOLI	ECULI	E TY	PE: I	ONA											
	(xi)	SEQ	JENCI	E DES	SCRII	PTIO	N: SI	EO II	O NO	:19:							
GGG	ATCCC							_									15
(2)	INFO	RMAT:	ION 1	FOR :	SEQ 3	ID NO	0:20	:									
	(i)	(A) (B) (C)	) LEI ) TYI ) STI	NGTH PE: 1 RAND	ARACT : 21 nucle EDNES GY: 1	base eic a SS: :	e pai acid sing:	irs									
	(ii)	MOLI	ECUL	E TY	PE: I	ONA											
	(xi)	SEQ	UENC	E DE	SCRII	PTIO	N: SI	EQ II	O NO	:20:							
GAA	rtcgg	CA CO	GAGC	CGAC	GG												21
(2)	INFO	RMAT:	ION :	FOR	SEQ :	ID N	0:21	:									
	(i)	(A (B (C	) LE ) TY ) ST	NGTH PE: 1 RAND	ARACT : 78 nucle EDNES	base eic a	e par acid sing	irs									
	(ii)	MOL	ECUL	Е ТҮ	PE: I	DNA											
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ II	D NO	:21:							
ATG	GAGCA	GA A	GCTG.	ATCT	C CG	AGGA	GGAC	CTG	CCCG	GGG (	CAGC	rggan	rc co	CAGO	CCAC	2	60
CCC	GCGCC	GG C	GGCC.	ATG													78
(2)	INFO	RMAT	ION	FOR	SEQ :	ID N	0:22	:									

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 26 amino acids

  - (B) TYPE: amino acid
    (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

Met Glu Gln Lys Leu Ile Ser Glu Glu Asp Leu Pro Gly Ala Ala Gly

Ser Ala Ala His Pro Ala Pro Ala Ala Met 20

- (2) INFORMATION FOR SEQ ID NO:23:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 33 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

GGATCCGCAG CCCACCCCGC GCCGGCGGCC ATG

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- (2) INFORMATION FOR SEQ ID NO:24:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 11 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:
- Gly Ser Ala Ala His Pro Ala Pro Ala Ala Met
- (2) INFORMATION FOR SEQ ID NO:25:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 21 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
      (D) TOPOLOGY: linear
  - (ii) MOLECULE TYPE: DNA

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	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
GGAC	CAAAGTG TGTGATGAAC C	21
(2)	INFORMATION FOR SEQ ID NO:26:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:	
CTCA	TCCTCC GTCTGATACT G	21
(2)	INFORMATION FOR SEQ ID NO:27:	
	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:	
GTAG	SATGTGG ATCAGCTTGG	20
(2)	INFORMATION FOR SEQ ID NO:28:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 19 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:	
AGGI	TGGAGAA TGGTCAAGG	19
(2)	INFORMATION FOR SEQ ID NO:29:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:	

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GTCATAGTCT GTCTCCTACT

(2)	INFORMATION FOR SEQ ID NO:30:	
	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:	
ACAT	AGACAG CGTGCCTACC	20
(2)	INFORMATION FOR SEQ ID NO:31:	
	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:	
TACA	ACCTTA GGGACACCAG	20
(2)	INFORMATION FOR SEQ ID NO:32:	
	<ul> <li>(i) SEQUENCE CHARACTERISTICS:</li> <li>(A) LENGTH: 20 base pairs</li> <li>(B) TYPE: nucleic acid</li> <li>(C) STRANDEDNESS: single</li> <li>(D) TOPOLOGY: linear</li> </ul>	
	(ii) MOLECULE TYPE: DNA	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
TGCT	TGAGCCT GCTCACGGTG	20
(2)	INFORMATION FOR SEQ ID NO:33:	
	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA	
7	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
CAA	GTGAACA GCACGTCC	18
(2)	INFORMATION FOR SEQ ID NO:34:	

(i) SEQUENCE CHARACTERISTICS:

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		<ul><li>(A) LENGTH: 21 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:34:	
GAC	ratct	CA AGGACCAGCT G	21
(2)	INFO	RMATION FOR SEQ ID NO:35:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:35:	
GGTT	CGGT	CC GAGCCCGG	18
(2)	INFO	RMATION FOR SEQ ID NO:36:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:36:	
GGAG	CGAT.	AC TCCAAGTAGG T	21
(2)	INFO	RMATION FOR SEQ ID NO:37:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:37:	

(2) INFORMATION FOR SEQ ID NO:38:

AGCGGGCCAG GCCCCTTC

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GCTCAGCTCC ACAAAGCGGC T

(2) INFORMATION FOR SEQ ID NO:42:

	(1)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:38:	
CATO	CTGG'	IC CAATGCGCTC	20
(2)	INFO	RMATION FOR SEQ ID NO:39:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
		SEQUENCE DESCRIPTION: SEQ ID NO:39:	
GCAC	CTGAG	GA AGTTAAACGA GC	22
(2)	INFO	RMATION FOR SEQ ID NO:40:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 22 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:40:	
		AA CTTCCTCAGT GC	22
(2)	INFO	RMATION FOR SEQ ID NO:41:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 21 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
e.	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:41:	

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(i) SEQUENCE CHARACTERISTICS:

		<ul><li>(A) LENGTH: 19 base pairs</li><li>(B) TYPE: nucleic acid</li><li>(C) STRANDEDNESS: single</li><li>(D) TOPOLOGY: linear</li></ul>	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:42:	
ACCA	AGCTC	CG CTCAGGTAG	19
(2)	INFO	RMATION FOR SEQ ID NO:43:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:43:	
TCC	AGGAG	CT GTGTGTTTGG	20
(2)	INFO	RMATION FOR SEQ ID NO:44:	
	(i)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 19 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:44:	
CCAC	STTTC	AC AGCGTGAGG	19
(2)	TNDO	DWARTON DOD GOO ED NO 45	
(2)		RMATION FOR SEQ ID NO:45:	
	(1)	SEQUENCE CHARACTERISTICS:  (A) LENGTH: 19 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
r	(ii)	MOLECULE TYPE: DNA	
	(xi)	SECUENCE DESCRIPTION: SEC ID NO.45.	

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CAGCATGAGG AGGAGGCAG

#### CLAIMS:

- 1. An isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding an amino acid sequence having homology to a regulator of gene expression or a derivative of said gene regulator.
- 2. An isolated nucleic acid molecule according to claim 1 wherein the regulator comprises a zinc finger domain of an (HC<sub>3</sub>)<sub>2</sub> type.
- 3. An isolated nucleic acid molecule according to claim 2 wherein the sequence of nucleotides or complementary sequence of nucleotides is selected from:
- (i) a nucleotide sequence set forth in SEQ ID NO:2;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:3;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridising under low stringency conditions to the nucleotide sequence set forth in (i), (ii) or (iii).
- 4. An isolated nucleic acid molecule according to claim 1 wherein said gene regulator is a guanine nucleotide exchange factor (GEF) or a derivative thereof.
- 5. An isolated nucleic acid molecule according to claim 4 wherein the sequence of nucleotides is selected from:
- (i) a nucleotide sequence set forth in SEQ ID NO:4 or 6;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:5 or 7;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridising under low stringency conditions to the

nucleotide sequence set forth in (i), (ii) or (iii).

- 6. An isolated nucleic acid molecule according to claim 1, wherein said gene regulator is a heat shock protein or is a heat shock binding protein or a derivative thereof.
- 7. An isolated nucleic acid molecule according to claim 6, wherein the sequence of nucleotides is selected from:
- (i) a nucleotide sequence set forth in SEQ ID NO:8;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:9;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridising under low stringency conditions to the nucleotide sequence set forth in (i), (ii) or (iii).
- 8. A genetic construct comprising a vector portion and a gene portion comprising a regulator of gene expression or a derivative thereof.
- 9. A genetic construct according to claim 8 wherein the gene portion comprises a zinc finger domain of (HC<sub>3</sub>)<sub>2</sub> type.
- 10. A genetic construct according to claim 9 wherein the gene portion comprises a nucleotide sequence selected from:
- (i) a nucleotide sequence set forth in SEQ ID NO:2;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:3;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridising under low stringency conditions to the nucleotide sequence set forth in (i), (ii) or (iii).

- 11. A genetic construct according to claim 8 wherein said gene portion is a nucleotide exchange factor (GEF) or derivative thereof.
- 12. A genetic construct according to claim 11 wherein the gene portion comprises a nucleotide sequence selected from:
- (i) a nucleotide sequence set forth in SEQ ID NO:4 or 6;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:5 or 7;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridising under low stringency conditions to the nucleotide sequence set forth in (i), (ii) or (iii).
- 13. A genetic construct according to claim 8 wherein the gene portion is a heat shock protein or a derivative thereof or a heat shock binding protein or derivative thereof.
- 14. A genetic construct according to claim 13 wherein the gene portion comprises a nucleotide sequence selected from:
- (i) a nucleotide sequence set forth in SEQ ID NO:8;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:9;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- (iv) a nucleotide sequence capable of hybridising under low stringency conditions to the nucleotide sequence set forth in (i), (ii) or (iii).
- 15. A nucleic acid molecule encoding a gene regulator having the identifying characteristics of a molecule selected from MCG4, MCG7 and MCG18 having respective amino acid sequences of SEQ ID NO:3, SEQ ID NO: 5 or 7 and SEQ ID NO:9.

- 16. A method of detecting a condition caused or facilitated by an aberration in mcg4, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg4 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.
- 17. A method of detecting a condition caused or facilitated by an aberration in mcg4, said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG4 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.
- 18. A method for detecting MCG4 or a derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG4 or its derivatives or homologues for a time and under conditions sufficient for an antibody-MCG4 complex to form, and then detecting said complex.
- 19. A method of detecting a condition caused or facilitated by an aberration in mcg7, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg7 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.
- 20. A method of detecting a condition caused or facilitated by an aberration in mcg7, said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG7 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.
- A method for detecting MCG7 or a derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG7 or its derivatives or homologues for a time and under conditions sufficient for an antibody-MCG7 complex to form, and then detecting said complex.

- 22. A method of detecting a condition caused or facilitated by an aberration in mcg18, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said mcg18 wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of said condition or a propensity to develop said condition.
- 23. A method of detecting a condition caused or facilitated by an aberration in *mcg18*, said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG18 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.
- 24. A method for detecting MCG18 or a derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG18 or its derivatives or homologues for a time and under conditions sufficient for an antibody-MCG18 complex to form, and then detecting said complex.

TCAG	TAAF	ACA (	AGAG	ACTO	G GG	ATCO				TGC Cys		53
AGA Arg												101
GAG Glu 25												149
CTG Leu												197
AAC Asn												245
CTC Leu												293
AAC Asn												341
			ACC Thr									389
			ACA Thr									437
			GTG Val 140									485
			TGG Trp									533
			AGC Ser									581
			GCT Ala			Gly						629
					Pro			Pro			GAT Asp	677

ACG Thr	cgg Arg	GAT Asp	GAT Asp 220	Asp	23G Arg	ACA Thr	CCA Pro	GGC Gly 225	CTC Leu	CAT H1s	GGA Gly	JAC Asp	T. Cys 230	GAC Asp	GAT · Asp	· 5:125"
								TTG Leu								773
AGG Arg	AGC Ser 250	CGG Arg	GCT Ala	GGG Gly	TCT Ser	CGG Arg 255	AAG Lys	CGG Arg	CCG Pro	CTG Leu	ACC Thr 260	CTG Leu	CTC Leu	CAG Gln	CGG Arg	821
GCG Ala 265	GGG Gly	CTG Leu	CTG Leu	CTA Leu	CTC Leu 270	TTG Leu	GGA Gly	CTG Leu	CTG Leu	GGC Gly 275	TTC Phe	CTG Leu	CCC Ala	CTC Leu	CTT Leu 280	869
								GCC Ala								917
								ATC Ile 305						TGA		962
GCC	CCT	rgc '	TTGT	GGCT	AG G	CCAG	CTA	G GA	rgtg	GGTT	CTG	rgga	GGA :	GAGG	CGGGGT	1022
AAT	GGG(2	AGG (	CTGA	GGC2	AC C	ICTT	CACTO	s cc	cctc	rccc	TCA	AGCC'	TAA (	GACA	CTAAGA	1082
CCC	CAGA	ccc i	AAAG	CCAA	GT C	CACC	AGAG:	r gg	CTCG	CAGG	CCA	GCC'	rgg :	AGTC	CCCGTG	1142
GGT	CAAG	CAT '	TTGT	CTTG	AC T	rgct	TTCT	2 000	GGT	CTCC	AGC	CTCC	GAC	CCCT	CGCCCC	1202
ATG.	AAGG:	AGC '	TGGC.	AGGT	GG A	AATA	AACA	A CA	ACTT'	TTAT						1242

3/32

### Figure 2

gb[AA155210]AA155210 mr98e01.rl Stratagene mouse embryonic carcinoma (#937317) Mus musculus cDNA clone 605496 5

Query: ! MGLCKCPKRKVTNLFCFEHRVNVCEHCLVANHAKCIVQSYLQWLQDSDYNFNCRLCNIPL 60

MGLCKCPKRKVTNLFCFEHRVNVCEHCLVANHAKCIVQSYLQWLQDSDYNFNCRLCN PL

Sbjet: 38 MGLCKCPKRKVTNLFCFEHRVNVCEHCLVANHAKCTVQSYLQWLQDSDYNPNCRLCNTPL 273

### Figure 3

dbj|D75913|CELK111G3F C.elegans cDNA clone ykl11g3 : 5' end, single read.

Query: 7 PKRKVTNLFCFEHRVNVCEHCLVANHAKCIVQSYLQWLQDSDYNPNCRLCNIPLASRETT 66 PKRKVTNLF +EHRVNVCE LV NH C+VQSYL WL D DY+PNC LC L +T

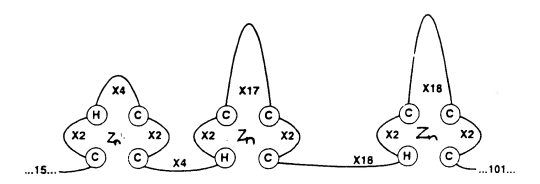
Sbjct: 1 PKRKVINLFXYEHRVNVCELXLVDNHPNCVVQSYLTWLTDQDYDPNCSLCKTTLXEGDTI 180

Query: 67 RLVCYDLFHWACLNERAAQLPRNTAPAGYQCP 98 98 PSCNGPIFPPNQ 109

RL C L HW C +E P TAP GY+CP P C+ +FPP+Q

Sbjct: 181 RLNCLHLLHWKCFDEWXGNFPDTTAPXGYRCP 276 275 PCCSQEVFPPDQ 310

Figure 4



### Figure 5

Query: 56 CNIPLASRETTRLVCYDLFHWACLNERAAQLPRNTAPAGYQCPSC 100 C+I L ++ + L C LF W C+ E A + + + + + CP C

Sbjct: 1222 CSICLENKNPSALFCGHLFCWTCIQEHAVAATSSASTSSARCPQC 1266

### Figure 6

gi|703468 (L29051) homologous to GATA-binding transcription factor [Schizosaccharomyces pombe]

Query: 35 CIVQSYLQWLQDSDYNPNCRLCNI 58

C + +W +D NP C C +

Sbjct: 175 CATTNTPKWRRDESGNPICNACGL 198

Query: 162 SSTPGPEEVDSASAAPAFYSQAPRPPASPGRPEQHTVIHMCNPEPLTHAPRKVYDTRDDD 221

+S PEE S S S P+ SP + +Q +I P +V + D

Sbjct: 441 ASILNPEEPPSNSDKQPSMSNGPKSEVSPSQSQQAPLIQSSTSPVSLQFPPEVQGSNVDK 500

Query: 222 RTPGLH 227 R L+

Sbjct: 501 RNYALN 506

Figure 7



gb|AA074703|AA074703 zm76g07.rl Stratagene neuroepithelium (#937231) Homo sapiens cDNA clone 531612 5' Length = 417

Plus Strand HSPs: Score = 818 (226.0 bits), Expect = 6.1e-103, Sum P(5) = 6.1e-103Identities = 206/259 (79%), Positives = 206/259 (79%), Strand = Plus / Plus 446 GGCCTCCCTCTGATCGATGAGGTGGTGAGCCCAGAGCCCGAGCCCCTCAACACGTCTGAC 505 49 GGGCTCCCTCTGATCGATGAGGTGATAAGCCCAGAGCCCGAGCCCCTCAATTCCTCAGAC 108 Sbict: 506 TYCTCTGACTGGTCTAGTTTTAATGCCAGCAGTACCCCTGGACCAGAGGAGGTAGACAGC 565 Query: Sbict: Query: 566 GCCTCTGCTGCCCCAGCCTTCTACAGCCAGGCCCCCCGGCCCCCAGCTTCCCCAGGCCGG 625 Sbict: Ouerv: 626 CCCGAGCACACACTGATCCACATGGGCAATCCTGAGCCCTTGACTCACGCCCCTAGG 685 Sbict: Query: 686 AAGGTGTATGATACGCGGG 704 Sbjct: Score = 230 (63.6 bits), Expect = 6.1e-103, Sum P(5) = 6.1e-103Identities = 50/55 (90%), Positives = 50/55 (90%), Strand = Plus / Plus Query: 398 GCACTGAGAGAGCTGGCCACAGTCAACTGGGCCCGGGCAGGACTGGGCCTCC 452 2 OCACTGAGAGAAAAGCTAGCCACAGTCAACTTGGCCCGGGCAGGACTGGGCTCCC 56 Score = 175 (48.4 bits), Expect = 6.1e-103, Sum P(5) = 6.1e-103Identities = 39/44 (88%), Positives = 39/44 (88%), Strand = Plus / Plus Query: 767 GCCTTGGGTTGGCTGGCCGGCTGCTAAGGAGCCGGGCTGGGTC 810 Sbict: 373 GCTCTGGGCTGGCCCAGCTGCTCAGGAGCCGGGCTGGGTC 416 Score = 139 (38.4 bits), Expect = 6.1e-103, Sum P(5) = 6.1e-103Identities = 31/35 (88%), Positives = 31/35 (88%), Strand = Plus / Plus Query: 731 GGAGACTGTGACGATGACAAGTACCGACGTCGGCC 765 336 GGAGACTSTGATGATGACAAATACCGCCGCCCGCC 370 Score = 133 (36.8 bits), Expect = 6.1e-103, Sum P(5) = 5.1e-103 Identities = 29/32 (90%), Positives = 29/32 (90%), Strand = Plus / Plus 701 CGGGATGATGACCGGACACCAGGCCTCCATGG 732 1411111111111111111 | 11111 | 11111 Sbjct: 305 CGGGATGATGACCGGACAGCAGCATTCATGG 336

#### Figure 8 continued

```
gb|AAl34788|AAl34788 zm8lg02.rl Stratagene neuroepithelium (#937231)
         Homo sapiens cDNA clone 532082 5'
         Length = 368
  Plus Strand HSPs:
 Score = 563 (155.6 \text{ bits}), Expect = 3.8e-87, Sum P(3) = 3.8e-87
 Identities = 147/190 (77%), Positives = 147/190 (77%), Strand = Plus / Plus
Query:
       498 CGTCTGACTTCTCTGACTGGTCTAGTTTTAATGCCAGCAGTACCCCTGGACCAGAGGAGG 557
          Sbjct:
       103 CCTCAGACTTCTCTGATTGGTCCAGCTTTAATGCCACCACCACCTCTGTGCAAGAGGAGA 162
       558 TAGACAGCGCCTCTGCTGCCCCAGCCTTCTACAGCCAGGCCCCCCGGCCCCCAGCTTCCC 617
Query:
       Sbjct:
       618 CAGGCCGGCCCGAGCAGCACACAGTGATCCACATGGGCAATCCTGAGCCCTTGACTCACG 677
Ouerv:
          Sbjct:
       223 CAAGCCGTCCCGAGCACACACACTCATACACATGGGGAGTACTGAAGCCCTGGCACACG 282
Query:
       678 CCCCTAGGAA 687
          1111 11111
Sbjct:
       283 CCCCAAGGAA 292
Score = 454 (125.4 bits), Expect = 3.8e-87, Sum P(3) = 3.8e-87
Identities = 94/98 (95%), Positives = 94/98 (95%), Strand = Plus / Plus
       Ouerv:
          Sbjct:
        Query:
       458 ATCGATGAGGTGGTGAGCCCAGAGCCCGAGCCCCTCAA 495
          Sbict:
       62 ATCGATGAGGTGATAAGCCCAGAGCCCGAGCCCCTCAA 99
Score = 219 (60.5 bits), Expect = 3.8e-87, Sum P(3) = 3.8e-87
Identities = 51/60 (85%), Positives = 51/60 (85%), Strand = Plus / Plus
Ouerv:
       702 GGGATGATGACCGGACACCAGGCCTCCATGGAGACTGTGACGATGACAAGTACCGACGTC 761
      309 GGATTGATGACCGGACAGCAGGCATTCATGGAGACTGTGATGATGACAAATACCGCCGCC 368
Sbjct:
```

### Figure 9

130	MCG4 MCG4 3.						PNCRLONI PL AGPVASALRE	
130	[ 229 ]						***x>	
NCG4			<del> </del>				****>	
1. 20 30 40 50 60 270 280 290 300 40 50 60 60 270 280 290 300 40 50 60 60 60 60 60 60 60 60 60 60 60 60 60		130	140	150	160	170	180	
372   30		KLATVIMARA						
2.								
3. 10 20 30 40 50 60    [ 229 ]				30	40	50	60	
229	[ 243 ]		·	_aqs*s*sip	*********	"tt"svq""r		
229	3	10	20	30	40	50	60	
5. 10 30 40 50 60 74 ] **********************************		********						
190   200   210   220   230   240	5.						60	
1.	[ 74]	*******	****smr**a	q**s*-sipq	tslig-pal-	mppp*lckrr	ep*lhlxlli>	
### MCG4 STAPPPASP GRPEQHTVIH MENPEPLTHA FRKVYDTRDD DRTPGLHGDC DDDKYRRPA  1.		R 190	200	210	220	230	240	
2.	MCG4	SPAPRPPASP	GRPEOHTVIH	MGNPEPLTHA	PRKVYDTRDD	DRTPGLHGDC	DODKYRRRPA	
2.					1			
2.			80					
3. 70 80 90 100 110 120 [ 229 ] gsp*sslpk* s*a-a*sht* gey*s*g*r- *kek*m*hg* **a*i********************************		V 70	80	90	pgp	"SINSWETVN	mtnt-aag1">	
3. 70 80 90 100 110 120  [ 229 ] gsp*sslpk* s*a-a*sht* gey*s*g*r- *kek*m*hg* ***a*i**** *******  4. 70 80 90 100 110 120  [ 86 ]p*sslpk* s*a-a*sht* gey*s*g*rp kesi*h*gmm tgqqafm*** ******c>		* /*****p**	s*******	**st*a*a**	_			
[ 229 ] gsp*sslpk* s*a-a*sht* gey*s*g*r- *kek*m*hg* ***a*i*** ***************************		V			1			
4. 70 80 90 100 110 120  [ 86 ]p*sslpk* s*a-a*sht* gey*s*g*rp kesi*h*gmm tgqqafm*** *******c> h  5. 70 80 90 100 110  [ 74 ] arl*allppq av*sstqsyt w*vlk*w-*t *qgk*m**** ***a*i**>								
[ 86 ]p*sslpk* s*a-a*sht* gey*s*g*rp kesi*h*gmm tgqqafm*** *******************  5.		gsp*sslpk*	s*a-a*sht*					
5. 70 80 90 100 110 [ 74 ] arl*allppq av*sstqsyt w*vlk*w-*t *qgk*m**** ***a*i**>  6.   100   [ 38 ]								
6.   100   250   260   270   280   290   300   MCG4   LGWIARILES RAGSEKERPLT LLQRAGILLL LGLIGFLALL ALMSRIGRAA ADSDENIJOPL 1.   130					h i			
6.   100   1	5.							
6.   100   1	[ 74 ]	arl*allppq	av*sstqsyt	w*vlk*w-*t	*qgk*m****	***a*i**>		
250 260 270 280 290 300  MCG4 LCWLARLLES RAGSRERPLT LLQRAGLILL LGLLGFLALL ALMSRLGRAA ADSDPNLDPL  1. 130					g			
250 260 270 280 290 300  MCG4 LCWLARLLES RAGSRERPLT LLQRAGLILL LGLLGFLALL ALMSRLGRAA ADSDPNLDPL  1. 130					100			
MCG4 LCWLARLLES RAGSRERPLT LLQRAGLILL LGLLGFLALL ALMSRLGRAA ADSDPNLDPL  1. 130	{ 38 }			*t	*q*******>	,		
1. 130		250	260	270	280	290	300	
{			RAGSRERPLT	LLQRAGLLLL	IGLIGFLALL	almsrigraa	ADSDPNLDPL	
	[ 372 ]	d	****>					
4. { 86 } s*-**>		s*-**>						
310		310						
MCG4 MINPHLIRVGPS	MCG4	MNPHIRVGPS		-				

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Figure 10 (Cont .ued)

Search Analysis for Sequence: MCG4

Search from 1 to 310

Date: September 22,1997

Matrix: pam250 matrix

Score Region from 1 to 310

Maximum possible score: 1598

#### Aligned sequences:

1. = EST AA074703 phase 1 translation

2. = EST AA134788 phase 3 translation

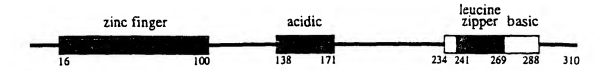
3. = EST AA134788 phase 2 translation

4. = EST AA074703 phase 3 translation

5. = EST AA074703 phase 2 translation

6. = EST AA134788 phase 1 translation

# FIGURE 11 Domains of MCG4



acidic domain consensus: 9/34 negatively charged amino acids, 0/34 positively charged

basic domain consensus: 13/55 positively charged amino acids, 0/55 negatively charged

leucine zipper domain consensus: LX<sub>6</sub>LX<sub>6</sub>RX<sub>6</sub>LX<sub>6</sub>L

alternate "novel" leucine zipper-like motif where leucine would not be aligned along the one surface of an alpha helix domain: (aa 261) LX<sub>6</sub>LXLX<sub>6</sub>L(aa 286)

FIGURE 12												
			Smallest									
			Sum									
		High	Probabili	ty								
Sequences producing !	High-scoring Segment Pairs:	Score	P(N)	N								
gnl PID e236178	(270752) F25B3.3 (Caenorhabditis ele	307	3.0e-124	8								
gi 1293099	(U53884) aimless RasGEF (Dictyosteli	202	7.8e-22	5								
gi 1655941	(U67326) Ras-GRF2 [Mus musculus]	152	3.6e-16	4								
pir  S30356	CDC25 protein homolog - yeast (Candi	150	2.2e-15	3								
	CELL DIVISION CONTROL PROTEIN 25	150	2.2e-15	3								
SP P28818 GNRP_RAT	GUANINE NUCLEOTIDE RELEASING PROTEIN	166	2.6e-15	3								
prf   1814463A	quanine nucleotide-releasing factor	166	2.6e-15	3								
pir   B46199	nucleotide-exchange-factor homolog c	167	1.le-14	1								
gn1 PID e238680	(X97560) hypothetical protein L1309	158	3.0e-14	3								
pir  S22693	CDC25 protein homolog - mouse /gi   50	167	3.7e-14	2								
	SCD25 PROTEIN /gi   457494 (M26647) SD	158	4.6e-14	3								
-,	STE6 PROTEIN /pir   S28098 ste6 prote	160	5.2e-14	2								
pir  S28407	CDC25 protein homolog - mouse	167	1.2e-13	3								
	GUANINE NUCLEOTIDE RELEASING PROTEIN	167	1.2e-13	3								
- :		153	2.0e-13	2								
gi 386047	(S62035) Ras-specific guanine nucleo	142	4.5e-13	2								
• • • • •	CELL DIVISION CONTROL PROTEIN 25 /pi	152	5.7e-13	3								
pir  S14177	SCD25 protein - yeast (Saccharomyces			3								
gi  433720	(L26584) CDC25 [Homo sapiens]	153	6.0e-13									
gn1 PID e241744	(Z68880) T14G10.2 (Caenorhabditis el	157	7.2e-13	1								
gi 3484	(X03579) CDC25 protein (aa 1-1588) [	136	3.4e-12	3								
T. I	CELL DIVISION CONTROL PROTEIN 25 /pi	136	3.4e-12	3								
gi 915328	(U24070) Muncl3-1 (Rattus norvegicus)	151	5.5e-12	1								
pir   A46199	nucleotide-exchange-factor homolog c	149	5.6e-12	1								
pdb   1PTR	Molecule: Protein Kinase C Delta Ty	136	1.5e-11	1								
gi 915330	(U24071) Munc13-2 [Rattus norvegicus]	150	1.6e-11	2								
gi 474982	(D21239) 'C3G protein' [Homo sapiens	131	3.3e-11	3								
gi 1763306	(U75361) Muncl3-3 [Rattus norvegicus]	153	6.4e-11	2								
gi 806957	guanine-nucleotide exchange factor C	128	7.8e-11	3								
sp Q03385 CNDS_MOUSE	GUANINE NUCLEOTIDE DISSOCIATION STIM	133	1.0e-10	2								
pir BVBYL1	LTE1 protein - yeast (Saccharomyces	139	1.9e-10	1								
gi  452242	(D21354) a putative guanine nucleoti	139	2.7e-10	1								
SP P07866 LTE1_YEAST	LOW TEMPERATURE ESSENTIAL PROTEIN /p	139	2.7e-10	1								
gi 509050	(Z22521) protein kinase C delta [Hom	137	4.0e-10	1								
gi 520587	(D10495) protein kinase C delta-type	137	4.6e-10	1								
SD P05130 KPC1_DROME	PROTEIN KINASE C, BRAIN ISOZYME (PKC	137	4.7e-10	1								
pir  S35704	protein kinase C (EC 2.7.1) delta	137	4.7e-10	1								
	PROTEIN KINASE C, DELTA TYPE (NPKC-D	137	4.7e-10	1								
pir  S40279	protein kinase C mu - human /pir   A5	137	4.9e-10	1								
sp P09215 KPCD_RAT	PROTEIN KINASE C. DELTA TYPE (NPKC-D	135	9.0e-10	1								
gi 520878	(234524) serine/threonine protein ki	133	1.8e-09	1								
gi 1519719	(U68142) RalGDS-like [Homo sapiens]	115	3.8e-09	3								
3-1	(and the same true frame and and											

## 12/32 FIGURE 13(a) (i)

MCG7 - Cloning of a novel human gene that encodes a guanine exchange factor

CGATTTCATTCCTCGCTCCCCACAGGTCCCTCTCCCCAAAATATTCCCATCTTGTCCTAG 60 I S F L A P H R S L S P K Y S H L V L CCCATCCCCAGACTATCTCAAGGACCAGCTGTCCCCACGCCCCCGACCTCCACTAGGCC 120 A H P P D Y L K D Q L S P R P R P P L G TGTGCCACCGGTGCCTGCAGGAAGACGCCCGGTCCCGGGCCGGGTTAGCCCCATGGGAA 180 LCHPLPAGRRPVPGRVSPMG T Q R L C G R G T Q G W P G S S E Q H V aggaggcgacctcgtccgcgggtttgcattctggggtggacgagctggGGGTTCGGTCCG 300 Q E A T S S A G L H S G V D E L G V R S E P G G R L P E R S L G P A H P A P A A TGGCAGGCACCCTGGACCTGGACAAGGGCTGCACGGTGGAGGAGCTGCTCCGCGGGTGCA 420 M A G T L D L D K G C T V E E L L R G C TCGAAGCCTTCGATGACTCCGGGAAGGTGCGGGACCCGCAGCTGGTGCGCATGTTCCTCA 480 I E A F D D S G K V R D P O L V R M F L TGATGCACCCTGGTACATCCCCTCCTCTCAGCTGGCGGCCAAGCTGCTCCACATCTACC 540 179 M M H P W Y I P S S Q L A A K L L H I Y AACAATCCCGGAAGGACAACTCCAATTCCCTGCAGGTGAAAACGTGCCACCTGGTCAGGT 600 Q Q S R K D N S N S L Q V K T C H L V R ACTGGATCTCCGCCTTCCCAGCGGAGTTTGACTTGAACCCGGAGTTGGCTGAGCAGATCA 660 WISAFPAEFDLNPELAEQI AGGAGCTGAAGGCTCTGCTAGACCAAGAAGGGAACCGACGGCACAGCAGCCTAATCGACA 720 K E L K A L L D Q E G N R R H S S L I D TAGACAGCGTCCCTACCTACAAGTGGAAGCGGCAGGTGACTCAGCGGAACCCTGTGGGAC 780 I D S V P T Y K W K R Q V T Q R N P V G AGAAAAAGCGCAAGATGTCCCTGTTGTTTGACCACCTGGAGCCCATGGAGCTGGCGGAGC 840 O K K R K M S L L F D H L E P M E L A E ATCTCACCTACTTGGAGTATCGCTCCTTCTGCAAGATCCTGTTTCAGGACTATCACAGTT 900 H L T Y L E Y R S F C K I L F Q D Y H S TCGTGACTCATGGCTGCACTGTGGACACCCCGTCCTGGAGCGGTTCATCTCCCTCTTCA 960 F V T H G C T V D N P V L E R F I S L F ACAGCGTCTCACAGTGGGTGCAGCTCATGATCCTCAGCAAACCCACAGCCCCGCAGCGGG 1020 N S V S O W V O L M I L S K P T A P Q R CCCTGGTCATCACACACTTTGTCCACGTGGCGGAGAAGCTGCTACAGCTGCAGAACTTCA 1080 A L V I T H F V H V A E K L L O L O N F ACACGCTGATGGCAGTGGTCGGGGGCCTGAGCCACAGCTCCATCTCCCGCCTCAAGGAGA 1140 NTLMAVVGGLSHSSISRLKE CCCACAGCCACGTTAGCCCTGAGACCATCAAGCTCTGGGAGGGTCTCACGGAACTAGTGA 1200 T H S H V S P E T I K L W E G L T E L V CGGCGACAGGCAACTATGGCAACTACCGGCGTCGGCTGCAGCCTGTGTGGGGCTTCCGCT 1260 T A T G N Y G N Y R R R L A A C V G F R TCCCGATCCTGGGTGTGCACCTCAAGGACCTGGTGGCCCTGCAGCTGGCACTGCCTGACT 1320 F P I L G V H L K D L V A L Q L A L P D GGCTGGACCCAGCCCGGACCCGGCTCAACGGGGCCAAGATGAAGCAGCTCTTTAGCATCC 1380 WLDPARTRLNGAKMKQLFSI TGGAGGAGCTGGCCATGGTGACCAGCCTGCGGCCACCAGTACAGGCCAACCCCGACCTGC 1440 LEELAMVTSLRPPVQANPDL TGAGCCTGCTCACGGTGTCTCTGGATCAGTATCAGACGGAGGATGAGCTGTACCAGCTGT 1500 LSLLTVSLDQYQTEDELYQL CCCTGCAGCGGGAGCCGCGCTCCAAGTCCTCGCCAACCAGCCCCACGAGTTGCACCCCAC 1560 SLQREPRSKSSPTSCT? CACCCCGGCCCCCGGTACTGGAGGAGTGGACCTCGGCTGCCAAACCCAAGCTGGATCAGG 1620 PPRPPVLEEWTSAAKPKLDQ CCCTCGTGGTGGAGCACATCGAGAAGATGGTGGAGTCTGTGTTCCGGAACTTTGACGTCG 1680

### FIGURE 13(a) (ii)

A L V V E H I E K M V E S V F R N F D V ATGGGGATGGCCACATCTCACAGGAAGAATTCCAGATCATCCGTGGGAACTTCCCTTACC 1740 D G D G H I S Q E E F Q I I R G N F P Y TCAGCGCCTTTGGGGACCTCGACCAGAACCAGGATGGCTGCATCAGCAGGGAGGAGGATGG 1800 L S A F G D L D Q N Q D G C I S R E B M TTTCCTATTTCCTGCGCTCCAGCTCTGTGTTGGGGGGGCGCATGGGCTTCGTACACAACT 1860 V S Y F L R S S S V L G G R M G F V H N TCCAGGAGAGCAACTCCTTGCGCCCGTCGCCTGCCGCCACTGCAAAGCCCTGATCCTGG 1920 P Q E S N S L R P V A C R H C K A L I L GCATCTACAAGCAGGGCCTCAAATGCCGAGCCTGTGGAGTGAACTGCCACAAGCAGTGCA 1980 G I Y K Q G L K C R A C G V N C H K Q C AGGATCGCCTGTCAGTTGAGTGTCGGCGCAGGGCCCAGAGTGTGAGCCTGGAGGGGTCTG 2040 K D R L S V E C R R R A Q S V S L E G S A P S P S P M H S H H H R A F S F S L P GCCCTGGCAGGCGAGGCTCCAGGCCTCCAGAGATCCGTGAGGAGGAGGTACAGACGGTGG 2160 R P G R R G S R P P E I R E E E V Q T V AGGATGGGGTGTTTGACATCCACTTGTAATAGATGCTGTGGTTGGATCAAGGACTCATTC 2220 BDGVFDIHL \* 5 CTGCCTTGGAGAAAATACTTCAACCAGAGCAGGGGGGCCTGGGGGTGTCGGGGCAGGAGGC 2280 TGGGGATGGGGTGGGATATGAGGGTGGCATGCAGCTGAGGGCAGGGCCAGGGCTGGTGT 2340 CCCTAAGGTTGTACAGACTCTTGTGAATATTTGTATTTTCCAGATGGAATAAAAAGGCCC 2400 GTGTAATTAACCTTC (A) n

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## FIGURE 13(b)

CGATTTCATTCCTCGCTCCCCACAGGTCCCTCTCCCCAAAATATTCCCATCTTGTCCTAG 60
CCCATCCCCCAGACTATCTCAAGGACCAGCTGTCCCCACGCCCCCGACCTCCACTAGGCC 120
TGTGCCACCCGCTGCCTGCAGGAAGACGCCCGGTCCCGGGCCGGGTTAGCCCCATGGGAA L80

p h g n

CGGGGTTCGGTCCGAGCCCGGTGGGAGGCTCCCGGAGCCTGGGCCCAGCCCACCC-240
g v r s e p g g r l p e r s l g p a h p

CGCGCCGGCGGCCATGGCAGGCACCCTGGACCAGGGCTGCACGGTGGAGGAGGT-360
a p a a M A G T L D L D K G C T V E E L

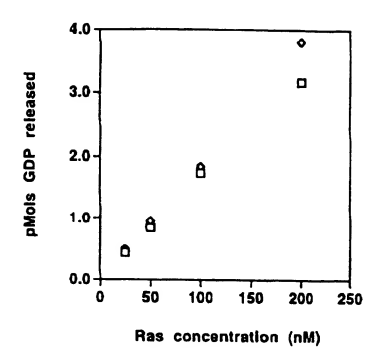
1	MAGTLDLDKGC:TVEELLRGCIEAFDDSGKVRDPQLVRMFLMMHPW	45
1	MSSKVEEDQHQELLTEDQLVARCVECFDVDEEDEVEDIEFVDALFLSHQW	50
46	YIPSQLAAKLLHIYQQSRKDNSNSLQVKTCHLVRYWISAFPAEFDLNPE	95
51	: : :     :   . :   :   . :   .   .	97
96	LAEQIKELKALLDQEGNRHSSLIDIDSVPTYKWKRQVTQRNPVGQKK	143
98	VCAQVVRLKTIAEDINENIRNGL.DVSALPSFAWLRAVSVRNPLAKQTIV	146
144	RKMSLLFDHLEPMELAEHLTYLEYR	168
147	RVDFETLPTPGTPPPFPIASKKFSLTAFSLSFVQASPSDISTSLSHIDYR	196
169	SFCKILFQDYHSFVTHGCTVDNPVLERFISLFNSVSQWVQLMILSKPTAP ::: :::: :::: :               :  :  :	218
197	VLSRISITELKQYVKDGHLRSCPMLERSISVFNNLSNWVQCMILNKTTPK	246
219	QRALVITHFYHYAEKLLOLONFNTLMAVVGGLSHSSISRLKETHSHVSPE :   ::      .:.    ::.  :   ::.  :   ::.	268
247	ERAEILVKFVHVAKHLRKINNFNTLMSVVGGITHSSVARLAKTYAVLSND	296
269	TIKLWEGLTELVTATGNYGNYRRRLAAC.VGFRFPILGVHLKDLVALQLA	317
297	IKKELTQLTNLLSAQHNFCEYRKALGACNKKFRIPIIGVHLKDLVAINCS	346
318	LPDWLDPARTRLNGAKMKQLFSILEELAMVTSLRPPV.QANPDLLSLLTV ::: : :       : .   .	366
347	GANFEKTKCISSDKLVKLSKLLSNFLVFNQKGHNLPEMNMDLINTLKV	394
367	SLDQYQTEDELYQLSLQREPRSKSSPTSPTSCTPPPRPPVLEEWTSAAKP	416
395		437
417	KLDQALVVEHIEKMVESVFRNFDVDGDGHISOEEFQIIRGNF}YLSAFGD	466
438	APDNATVSKHISAMVDAVFKHY <u>DHDRDGFISOEEFQ</u> LIAGNFPFIDAFVN	487
467	LDONODGCISREEMVSYFLRSS.SVLGGRMGFVHNFOESNSLRPVACRHC	515
488	IDVDMDGOISKDELKTYFMAANKNTKDLRRGFKHNFHETTFLTPTTCNHC	537
516	KALILGIYKOGLKCRACGVNCHKOCKDRLSVECRRRAQSVSLEGSAPSPS	565
538	NKLLWGILROGFKCKDCGLAVHSCCKSNAVAECRRKSSSNLTRAAEWFAS	587
566	PMHSHHHRAFSFSLPRPGRRGSRPPEIREEEVQTVEDGVFDIHL 609	
588		

human	CGATTICATT CCT	reservece e	CACAGGTCCC	TCTCCCCAAA	ATATTCCCAT	CITGTCCTAG 60
human	CCCATCCCCC AGA	ACTATOTO A	AAGGACCAGC	TGTCCCCACG	CCCCCGACCT	CCACTAGGCC 120
human	TGTGCCACCC GCT	receteca e	GAAGACGCC	CCCTCCCCCC	CCGGGTTAGC	CCCATGGGAA 180
human	CGCAGCGCCT GTC	TGGCCGC (	GGACTCAAG	GCTGGCCTGG	CTCAAGTGAA	CAGCACGTCC 240
mouse			***tcag**	****ag****	f*******	***a*g***t>
human	AGGAGGCGAC CTC	CTCCCCCC C	GTTTGCATT	CTGGGGTGGA	CGAGCTCGGG	GTTCGGTCCG 300
					acago	•
					1	
					1	
mouse	g*****t**a **-				-	
human	AGCCCGGTGG GAG	SCICCCC C	BAGCGCAGCC	TGGGCCCAGC	CCACCCCGCG	CCGGCGGCCA 360
mouse	***a*t**** ***	****tga *	***t*t*a*t	*****	***-*tg**a	****a****>
human	TOGCAGGCAC CCT	rggacetg c	SACAAGGGCT	GCACGGTGGA	GGAGCTGCTC	CGCGGGTGCA 420
mouse	****ga**** t**	******	*******	****C****	******	**t**c**t*>
human	TCGAAGCCTT CGA	ATGACTCC G	GGAAGGTGC	GGGACCCGCA	GCTGGTGCGC	ATGTTCCTCA 480
mouse	******** ***	******	*a*****	*a**t**a**	***a*****	******
human	TGATGCACCC CTC	STACATO C	CCTCCTCTC	AGCTGGCGGC	CAAGCTGCTC	CACATCTACC 540
mouse	******** ***	******a *	*******	******tt*	g**a*****	***t****t*>
human	AACAATCCCG GAA	AGGACAAC T	CCAATTCCC	TGCAGGTGAA	AACGTGCCAC	CTGGTCAGGT 600
mouse	*g******	*****	********	*a***a****	******	C*******
human	ACTGGATCTC CGC	CTTCCCA C	CGGAGTTIG	ACTIGAACCC	GGAGTTGGCT	GAGCAGATCA 660
mouse	**** a**	*****	"*a****c*	********	a***c****	**a******
human	AGGAGCTGAA GGC	TCTGCTA C	BACCAAGAAG	GGAACCGACG	GCACAGCAGC	CTAATCGACA 720
mouse	*********	*****	*******	******ca*	********	**c******
human	TAGACAGCGT					730
mouse	*c**g**t**					

# 17/32

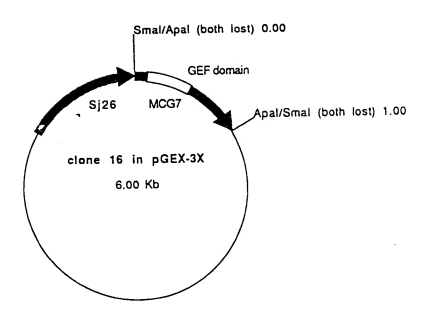
CA	'CGC	CTC	GGA	LAGO	GAC	GT.	TTGG	GG	CCC	STGC	TT	CAC	AGT	GAC	TG	CTC	TGA	VAGC	CAAA	60
TG	GTC	:GGA	AAC	CGT	CTAC	CCC	CTC	TC	CTAC	GC(	CGC	CTA	GTG	GGG	ACC	CC	ACC	GCC	TGCG	120
									*	A	R	L	V	G	T	P	T	A	C>	
GC	TGC	CCC	TCC	CA	\GT	וככי	rccc	TG	TGC	SCC#	\GG(	ATC	CAG	GTC	TCC	AG1	CTC	CGA	GCTG	180
G	C	P	S	Q	V	P	P	C	W	₽	G	I	Q	V	S	S	L	R	A>	
CG	GAG	AAC	CCA	CCC	CC	CAT	rGCG	GC7	GCC	בכם	TTC	CAT	TCG	ACC	CTG	TGG	GGA	GCC	AGGC	240
A	E	N	P	P	P	H	A	Α	A	P	F	H	s	T	L	W	G	A	R>	
TT	CCG	GGG	CCC	CGI	TCC	TCC	TGT	GTO	AAC	TGG	GCC	CCC	CGC	CCC	CAT	TCC	CAG	ACA	TCAA	300
L	₽	G	P	R	S	s	C	V	N	W	A	P	R	P	H	S	Q	T	S>	
GG	CCG	CGT	CTC	CAG	ATA	\GC(	ACG	ATI	TCA	TTC	CTC	GCT	CCC	CAC	AGG	TCC	CTC	TCC	CCAA	360
R	P	R	L	Q	I	A	T	I	S	P	L	A	P	H	R	s	L	s	P>	
AA	TAT	TCC	CAT	CTI	GTC	CTA	GCC	CAT	CC.	CCA	GAC	TAT	CTC	AAG	GAC	CAG	CTG	TCC	CCAC	420
K	Y	s	H	L	V	L	A	H	P	P	D	Y	L	K	D	Q	L	s	P>	
GC	CCC	CGA	CCT	CCA	CTF	\GGC	CTG	TGC	CAC	CCG	CTG	CCT	GCA	GGA	AGA	.CGC	CCG	GTC	CCGG	480
R	P	R	P	P	L	G	L	C	H	P	L	P	A	G	R	R	₽	V	P>	
GC	CGG	GTT	AGC	CCC	ATC	GGA	ACG	cag	cgc	ctg	tgt	ggc	cgc	999	act	caa	ggc	tgg	cctg	540
				P		_	n													
G												G							P>	
gc	tca	agt	gaa	cag	cac	gto	cag	gag	gcg	acc	tcg	tcc	gcg	ggt	ttg	cat	tct	<b>3</b> 33	gtgg	600
				_			_					S								
ac	gag	ctg	gGG	GTT	CGC	TCC													CCAG	660
D	-	L	_			_	-	_	_	_	-	L	_	_		_	L	-	_	
CC	CAC	CCC	GCG	CCG	GCG	GCC	' <u>ATG</u>	GCA	'GGC	ACC	CTG	GAC	CTG	GAC	AAG	GGC	TGC	ACG	GTGG	720
Δ	H	P	A	Þ	Α	A	м	A	G	T	T.	D	T.	D	K	G	C	T	٧>	

FIGURE 17



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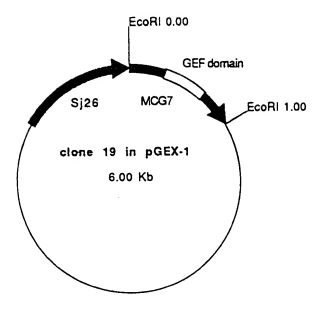
19/32
FIGURE 18 (Cont. I)



Plasmid name: clone 16 in pGEX-3X

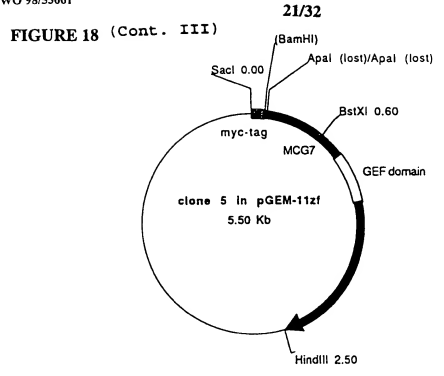
Plasmid size: 6.00 kb

FIGURE 18 (Cont. II)



Plasmid name: clone 19 in pGEX-1

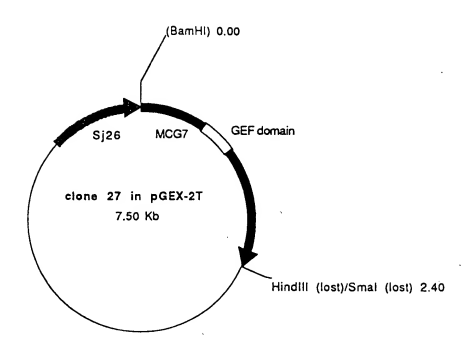
Plasmid size: 6.00 kb



Plasmid name: clone 5 in pGEM-11zf

Plasmid size: 5.50 kb

FIGURE 18 (Cont. IV)



Plasmid name: clone 27 in pGEX-2T

Plasmid size: 7.50 kb

# FIGURE 19

GCC	.GCC	GCC	Met 1													49
												Ala			CGG	97
			AGT Ser													145
			GAA Glu													193
			CGG Arg 65													241
			GAG Glu							_						289
_			GAC Asp				_	-								337
			CAT His													385
			GCA Ala													433
															CTG Leu	481
-			Leu							_			His		ATT Ile	529
												Met			AAG Lys	
GAT	CGG	ATO	ATC	ACA	GCC	TTC	TAC	AAC	GAA	GCC	CGG	GCA	cae	GCC	AGG	625

WO 98/53061 FIGURE 19 (cont.   led)					24/32					PCT/AU98/00380						
Asp 190		Ile	Ile	Thr	Ala 195	Phe	Tyr	Asn	Glu	Ala 200	Arg	Ala	Arg	Ala	Arg 205	
GCC	AAC	AGA	GGC	ATC	CTT	CAG	CAG	GAG	CGA	CAA	CGG	CTA	GGG	CAG	CGG	673
Ala	?. sn	Arg	Gly	Ile 210	Leu	Gln	Gla	Glu	Arg 215	Gln	Arg	Leu	Gly	Gln 220	Arg	
CAG	CCG	CCA	CCA	TCC	GAG	CCA	ACC	CAA	GGC	CCC	GAG	ATC	GTG	CCC	CGG	721
Gln	Pro	Pro	Pro 225	Ser	Glu	Pro	Thr	Gln 230	Gly	Pro	Glu	Ile	Val 235	Pro	Arg	
			CCC Pro		GGGG	CTC	ACCT	GGAT	GG G	GCCT	GCAG	T GC	GTTC	CCGC		773

74

ć.

:

Length = 376

Score = 138 (63.7 bits), Expect = 1.2e-10, P = 1.2e-10
Identities = 25/62 (40%), Positives = 39/62 (62%)

Query: 35 YYELLGVHPGASTEEVKRAFFSKSKELHPDRDPGNPSLHSRFVELSEAYRVLSREQSRRS 94

YYE+LGV A E+++A+ + + HPDR+ G+ ++F E+ EAY VL+ Q R +

Sbjct: 6 YYEILGVSKTAEEREIRKAYKRLAMKYHPDRNQGDKEAEAKFKEIKEAYEVLTDSQKRAA 65

Query: 95 YD 96 YD Sbjct: 66 YD 67

>gi|1703590 (U80439) contains similarity to a DNAJ-like domain (Caenorhabditis elegans) Length = 345

Score = 98 (45.2 bits), Expect = 5.2e-12, Sum P(3) = 5.2e-12 Identities = 17/37 (45%), Positives = 28/37 (75%)

Query: 28 QRSRPSTYYELLGVHPGASTEEVKRAFFSKSKELHPD 64

++ R T+YE+LGV A+ E+K AF+++SK++HPD

Sbjct: 22 KKIRQRTHYEVLGVESTATLSEIKSAFYAQSKKVHPD 58

Score = 74 (34.1 bits), Expect = 5.2e-12, Sum P(3) = 5.2e-12 Identities = 17/32 (53%), Positives = 19/32 (59%)

Ouery: 71 SLHSRFVELSEAYRVLSREOSRRSYDDQLRSG 102

S + F+EL AY VL R RR YD QLR G

Sbjct: 64 SATASFLELKNAYDVLRRPADRRLYDYQLRGG 95

Score = 39 (18.0 bits), Expect = 5.2e-12, Sum P(3) = 5.2e-12 Identities = 10/42 (23%), Positives = 19/42 (45%)

Query: 162 LLMLAGMGLHYIAFRKVKQMHLNFMDEKDRIITAFYNEARAR 203

L+++AG Y+ Q L+ + ++D I F + R

Sbjct: 158 LVLVAGYNGGYLYLLAYNQKQLDKLIDEDEIAKCFLRQKEFR 199

>gn1|PID|e281266 (Z81030) C01G10.12 (Caenorhabditis elegans)
Length = 191

Score = 96 (44.3 bits), Expect = 1.8e-09, Sum P(3) = 1.8e-09 Identities = 17/41 (41%), Positives = 27/41 (65%)

Query: 35 YYELLGVHPGASTEEVKRAFFSKSKELHPDRDPGNPSLHSR 75

YYE++GV A+ +E++ AF K+K+LHPD+ + SR

Sbjct: 19 YYEIIGVSASATRQEIRDAFLKKTKQLHPDQSRKSSKSDSR 59

Score = 54 (24.9 bits), Expect = 1.8e-09, Sum P(3) = 1.8e-09 Identities = 10/22 (45%), Positives = 15/22 (68%)

Query: 75 RFVZLSEAYRVLSREQSPFSYD 96

+F+ + EAY VL E+ R+ YD

Sbjct: 71 QFMLVKEAYDVLRMEEKRKEYD 92

Score = 35 (16.1 bits), Expect = 1.8e-09, Sum P(3) = 1.8e-09 Identities = 9/44 (20%), Positives = 22/44 (50%)

Query: 141 QGPQLRQQQHKQNKQVLGYCLLLMLAGMGLHYIAFRKVKQMHLN 184

+ P+ + KQ ++L ++A +G + + RK++ L+

Sbjct: 145 RNPEDEYLREKOKNRMLVVLAATVMALIGANTVYIRKLQADRLS 188

>sp[Q10209]YAY1\_SCHPO HYPOTHETICAL 44.8 KD PROTEIN C4H3.01 IN CHROMOSOME I >gi|1184014 (269380) unknown [Schizosaccharomyces pombe] Length = 392Score = 84 (38.8 bits), Expect = 4.1e-08, Sum P(3) = 4.1e-08Identities = 13/36 (36%), Positives = 25/36 (69%) Ouerv: 35 YYELLGVHPGASTEEVKRAFFSKSKELHPDRDPGNP 70 YY+LLG+ A+ ++K+A+ + + HPD++P +P 9 YYDLLGISTDATAVDIKKAYRKLAVKYHPDKNPDDP 44 Sbict: Score = 64 (29.5 bits), Expect = 4.1e-08, Sum P(3) = 4.1e-08Identities = 14/40 (35%), Positives = 23/40 (57%) 75 RFVELSEAYRVLSREQSRRSYDDQLRSGSPPKSPPTTVHD 114 Ouerv: +F ++SEAY+VL E+ R YD + + P+ T +D Sbjct: 50 KFQKISEAYQVLGDEKLRSQYDQFGKEKAVPEQGFTDAYD 89 Score = 37 (17.1 bits), Expect = 4.1e-08, Sum P(3) = 4.1e-08Identities = 9/29 (31%), Positives = 15/29 (51%) Ouerv: 190 DRIITAFYNEARARANRGILQQERQRL 218 DR A E A A+ + +++ RQR+ 149 DRKKNAQIREREALAKREQEMIEDRRQRI 177 Sbict: Score = 33 (15.2 bits). Expect = 0.00081, Sum P(3) = 0.00081

Query: 140 PQGPQLRQQQHKQNKQVLG 158 PQG + Q+ + QVLG

Identities = 8/19 (42%), Positives = 11/19 (57%)

Sbjct: 44 PQGASEKFQKISEAYQVLG 62

#### FIGURE 23

Score  $\approx$  153 (70.6 bits), Expect = 9.7e-13, P = 9.7e-13 Identities = 27/71 (38%), Positives = 44/71 (61%)

Query: 26 AGQRSRPSTYYELLGVHPGASTEEVKRAFFSKSKELHPDRDPGNPSLHSRFVELSEAYRV 85
+ R + YY LGV A+ +++K+A++ +K+ HPD + +P +F ++SEAY V

Sbjct: 72 SSSRMQAKDYYATLGVAKNANAKDIKKAYYELAKKYHPDINKDDPDASKKFQDVSEAYEV 131

Query: 86 LSREQSRRSYD 96 LS +Q RR YD Sbjct: 132 LSDDQKRREYD 142

MCG18 HDJ-2 HDJ-1 HSJ1	SRLLGAA MVKETTYYDVLGVKPNATQEELKKAYRKLALKYHPDKNPNEGEXFKQISQAYEV MGKDYYQTLGLARGASDEEIKRAYRRQALRYHPDKNKEPGAEEKFKEIAEAYDV M-ASYYEILDVPRSASADDIKKAYRRKALQWHPDKNPDNKEFAEKKFKEVAEAYEV
MCG18 HDJ-2 HDJ-1 HSJ1	ACQRSRPSTYYELLGVHPGAST-EEVKRAFFS LSDAKKRELYDKGGEQAIKEGGAGGGFGSPMDIFDMFFGGG LSDPRKREIFDRYGEBGLKGSGPSGGSGGGANGTSFSYTFHGDPHAMFAEFFG LSDKHKREIYDRYGRBGLTGTGTGPSRAEAGSGGPGFTFT-FKSPEEVFREFFG
MCG18 HDJ-2 HDJ-1 HSJ1	KSKELHPDRDPGNPSLHSRFVELSEAYRVLSREQSRRSYDDQLRSGSPPKSPRT GRMQRERRGKNVVHQLSVTLEDLYNGATRKLALQKNVICDKCEGRGGKKGAVECCPNCRG GRNPFDTFFGQRNGEEGMDIDDPFSGFPMGMGGFTNVNFGRSRSAQEPARKQDPPVT SGDPFAELFDDLGPFSELQNRGSRHSGPFFTFSSSFPGHSDFSSSSFSFSPGAGAFRS
MCG18 HDJ-2 HDJ-1 HSJ1	TVHDKSAHQTHSSWTPPNAQYWSQFHSVRPQGPQLRQQQHKQN TGMQIRIHQIGPCMVQQIQSVCMECQGHGERISPK-DRCKSCNGRKIVREKKILEVHIDK HDLRVSLEEIYSGCTKKMKISH-KRLNPDGKSIRNEDKILTIEVKK VSTSTTFVQGRRITTRRIMENGQ-ERVEVEEDGQLKSVTINGVPD
MCG18 HDJ-2 HDJ-1 HSJ1	KQVLGYCLLLMLAGMGLHYIAFRKVKQMHLNFMDE-KDRIITAFYNEARARARAN GMKDGQKITFHGEGDQEPGLEPGDIIIVLDQKDHAVFTRRGEDLFMCMDIQLVEALCGFQ GMKEGTKITFPKEGDQTSNNIPADIVFVLKDKPHNIFKRDGSDVIYPARISLREALCGCT DLARGLELSR-REQQP-SVTSRSGGTQVQQTPASCPLD-SDLSEDEDLQLAMAYSLSE
MCG18 HDJ-2 HDJ-1 HSJ1	RGILQQERQRLCQRQPP-PSEPTQGPEIVPRGAGP
MCG18 HDJ-2 HDJ-1 HSJ1	SPDKLSLLEKLLPERKEVEETDEMDQVELVDFDPNQERRRHYNGEAYEDDEHHPRGGVQC PQTSRTVLEQVLPILIQILTGGSDSLWEEKRGVS
MCG18 HDJ-2 HDJ-1 HSJ1	ots

<sup>\* =</sup> amino acid identity in all 4 proteins

<sup>. =</sup> conservative substitution

CA	AGG	AGC	CTC	TGC	CTC	CCC	GTC	GTC	GTC	ATG	CCG	TCC	CTC	TTC	CTC	CAG	CTG	CCC	CTGC	60
										M	P	s	L	L	L	Q	L	P	L	10
GC	СТА	TGC	CGG	CTC	TGG	CCG	CAT	'AGC	CTI	TCC	ATC	CGA	.CTI	CTC	ACA	.GCC	GCC	ACA	.GGGC	120
R	L	C	R	L	W	P	Н	s	L	s	I	R	L	L	T	A	A	τ	G	30
AG	CGG	TCT	GTC	CCI	'ACT	'AAT	TAC	TAT	'GAA	TTG	TTG	GGC	GTG	CAT	CCG	GGT	GCC	AGC	GCTG	180
Q	R	s	v	P	T	N	Y	Y	Ε	L	L	G	V	Н	P	G	A	s	A	50
AA	GAG.	ATT	AAA	CGT	GCI	TTT	TTC	ACC	AAG	TCA	AAA	GAG	CTA	CAC	CCT	GAT	CGA	GAC	CCTG	240
E	E	I	ĸ	R	A	F	F	T	K	·s	ĸ	E	L	Н	P	D	R	D	P	70
GG.	AAC	CCA	GCC	CTG	CAT	AGC	CGC	TTI	GTG	GAG	CTG	TAA	GAG	GCA	TAT	CGA	GTG	CTC	AGTC	300
G	N	P	A	L	н	s	R	F	v	E	L	N	E	Æ	Y	R	v	L	s	90
GTY	GAG	GAA	AGT	CGT	CGT	AAC	ТАТ	GAC	CAC	CAG	CTG	CAT	TCA	GCC	AGT	CCT	CCA	AAG	TCTT	360
R	E	E	s	R	R	N	Y	D	н	Q	L	Н	\$	A	s	P	P	K	s	110
CA	GGG	AGC.	ACA	GCC	GAG	CCT	AAG	TAT	'ACG	CAA	CAG	ACA	CAC	AGC	AGC	TCC	TGG	GAA	cccc	420
s	G	s	T	A	E	P	K	Y	T	Q	Q	T	Н	s	S	s	W	E	P	130
CC	AAC	GCT	CAA	TAC	TGG	GCC	CAG	TTC	CAC	AGT	GTG	AGG	CCG	CAG	GGG	CCG	GAG	TCA	AGGA	480
P	N	A	Q	. Y	W	A	Q	F	н	s	v	R	P	Q	G	P	Ε	s	R	150
λG	CAG	CAG	CGT	AAA	CAC	AAC	CAG	CGG	GTC	CIG	GGG	TAC	TGC	CTC	CIG	CTC.	ATG	GTG	GCAG	540
K	Q	Q	R	K	Н	N	Q	R	V	L	G	Y	С	L	L	L	M	V	A	170
GC.	ATG	GGC	CTG	CAC	TAT	GTT	GCC	TTC	AGG	AAG	CTG	GAG	CAG	GTG	CAT	CGC	AGC	TTC.	ATGG	600
G	M	G	L	Н	Y	V	A	F	R	ĸ	L	E	Q	V	H	R	s	F	M	190
AΤ	GAA	AAG	GAC	CGG	ATC	ATT	ACA	GCC	ATC	TAC	TAA	GAC	ACT	CGG	GCC	AGG	GCC	AGG	GCCA	660
D	E	ĸ	D	R	r	I	T	A	I	Y	14	D	T	R	A	R	A	R	A	210
AC.	AG.N	GCC	AGG	ATI	CAG	CAG	GAG	CGC	CAC	GAG	AGG	CAG	CAG	CCT	CGG	GCA	GAA	ccc	TCCC	720
N	R	A	R	I	Q	Q	E	R	Н	E	R	Q	Q	P	R	A	E	P	s	230
TG	CCT	CCA	G <b>A</b> A	AGC	TCC	AGG	ATC	YTA:	ccc	CAG	GAC	ACA	AGC	CCC	TGA	.GAG	GCT	TAA	CTAA	780
L	P	P	E	s	s	R	I	M	P	Q	D	T	s	P	•					245
ΑT	CGG	ACC	TTC	AT	rgg1	rcci	CTC	cci	rgci	rgcc	TGI	CCA	(GA)	CT	CAC	GTC	CAA	<b>AAT</b>	ACTC	840
AT	TT	CAC	(A)	n																849

# FIGURE 26

human MCG18 mouse MCG18	MPPLLPLRLCRLWPRNPPSRLLGAAAGQRSRPSTYYELLGVHPGASTEEVKRAFFSK MPSLLLQLPLRLCRLWPHSLSIRLLTAATGQRSVPTNYYELLGVHPGASAEEIKRAFFTK
human MCG18	SKELHPDRDPGNPSLHSRFVELSEAYRVLSREQSRRSYDDQLRSGSPPKSPRTTVHDKSA
mouse MCG18	SKELHPDRDPGNPALHSRFVELNEAYRVLSREESRRNYDHQLHSASPPKSSGSTAEPRYT
human MCG18	HOTHSS-WTPPNAQYWSOFHSVRPQGPQLRQQQHKQNKQVLGYCLLLMLAGMGLHYLAFR
mouse MCG18	QQTHSSSWEPPNAQYWAQFHSVRPQGPESRKQQRKHNQRVLGYCLLLMVAGMGLHYVAFR
human MCG18	KVKQMHLNFMDEXDRIITAFYNEARARARANRGILQQERQRLGQRQPPPSEPTQGPE
mouse MCG18	KLEQVHRSFMDEXDRIITATYNDTRARARANRARIQQERHERQQPRAEPSLPPESSR
human MCG18	IVPRGAGP
mouse MCG18	IMPQDTSP

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### FIGURE 27

ttgaagtctagccccatcctggtccaatgcgctcttggtagcctcctttcccagctgccc 60
\* S L A P S W S N A L L V A S F P S C P

p A A M P P L L P L R L C R L W P R N P>

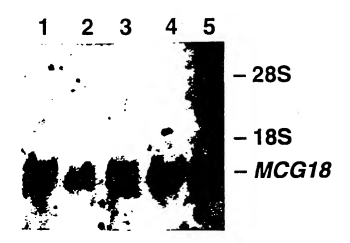


FIGURE 28

### INTERNATIONAL SEARCH REPORT

International Application No.

PCT/AU 98/00380

Α.	CLASSIFICATION OF SUBJECT MATTER							
Int Cl <sup>6</sup> :	C12N 15/12; C07K 14/47; C07K 16/18; G01N 3	3/53						
According to	International Patent Classification (IPC) or to both	national classification and IPC						
	FIELDS SEARCHED							
	Minimum documentation searched (classification system followed by classification symbols)  UC: WPAT (D gene) Sequences provided by Applicant							
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched								
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) :EMBL, Genebank, Swiss Prot and PIR: Sequences provided by applicant :								
C.	DOCUMENTS CONSIDERED TO BE RELEVANT							
Category*	Citation of document, with indication, where app	propriate, of the relevant passages	Relevant to claim No.					
P,X	Kedra D, Seroussi E, Fransson I, Trifunovic Blennow E, Mehlin H, Dumanski J, Human C 611-619 The germinal centre kinase gene and located in the vicinity of the PYGM gene on I EMBL AC Y12339  Guru S C, Agarwal S K, Manickain P, Olufe July 1997 7(7) 725-735. A transcript map for the multiple endocrine neoplasia type I locus	Genetics, October 1997 100(5-6) d a novel CDC25-like gene are 11q13  Temi S E, et al Genome Research, for the 2.8-Mb region containing 11.4-5, 8, 11-12, 11.4-5, 11.4-						
x	TREMBL AC 014616  Further documents are listed in the continuation of Box C	See patent family an	лех					
"A" document of color	al categories of cited documents:  ment defining the general state of the art which is onsidered to be of particular relevance or document but published on or after the national filing date ment which may throw doubts on priority claim(s) such is cited to establish the publication date of ere citation or other special reason (as specified) ment referring to an oral disclosure, use, bition or other means ment published prior to the international filing but later than the priority date claimed	priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art						
Date of the ac	tual completion of the international search	Date of mailing of the international sear	<del>-</del>					
16 July 1998	iling addragg of the ISA/AII	2 0 JUL 1998	<u> </u>					
AUSTRALIA PO BOX 200 WODEN AC AUSTRALIA		GILLIAN ALLEN Telephone No.: (02) 6283 2266	7(7(1600)					

# INTERNATIONAL SEARCH REPORT

International Application No.
PCT/AU 98/00380

Box 1	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This Inter	national Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:
2.	X Claims Nos.: 1, 2, 4, 6  because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:  They are to known groups of proteins and lack distinguishing features which would enable a meaningful search.
3.	Claims Nos.:  because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a)
Вох П	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	rnational Searching Authority found multiple inventions in this international application, as follows:
zinc fi Invent which Invent	tion 1, defined by claims 2, 3, 9, 10, 16-18, is to nucleotide sequences, amino acid sequences and proteins with a inger domain.  ion 2, defined by claims 4, 5, 11, 12, 19-21, is to nucleotide sequences and amino acid sequences and proteins are guanine exchange factors.  ion 3, defined by claims 6, 7, 13, 14, 22-24, is to nucleotide sequences and amino acid sequences and proteins are heat shock proteins or heat shock binding proteins.
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest  The additional search fees were accompanied by the applicant's protest.  X  No protest accompanied the payment of additional search fees.

### INTERNATIONAL SEARCH REPORT

international Application No.

PCT/AU 98/00380

C (Continua	tion) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
P,X	EMBL AC AF012106 DT 6 November 1997 Lloyd S E and Thakker R V DE Homo Sapiens DnaJ protein (HSPF <sub>2</sub> )mRNA, complete cds	1,6-8,13- 15,22-24
P,X	EMBL AC AF 036875 DT 20 May 1998 Silins G, Grimmond S, Hayward N DE Mus musculus multiple endocrine neoplasia type I candidate protein number 18 mRNA, complete cds	1,6-8,13- 15,22-24
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